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(54) Title: DETECTION OF FUNGAL PATHOGENS US	SING T	HE POLYMERASE CHAIN REACTION

DNA sequences from the Internal Transcribed Spacer of the ribosomal RNA gene region are described for different species and strains of Septoria, Pseudocercosporella, Fusarium and Mycosphaerella. Specific primers from within these sequences are identified as being useful for the identification of the fungal isolates using PCR-based techniques.

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# DETECTION OF FUNGAL PATHOGENS USING THE POLYMERASE CHAIN REACTION

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#### FIELD OF THE INVENTION

The present invention relates to the use of species-specific primers in polymerase chain reaction assays for the detection of fungal pathogens. The use of these primers enables the detection of specific isolates of fungal pathogens and the monitoring of disease development in plant populations.

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#### **BACKGROUND OF THE INVENTION**

Diseases in plants cause considerable crop loss from year to year resulting both in economic deprivation to farmers and additionally in many parts of the world to shortfalls in the nutritional provision for local populations. The widespread use of fungicides has provided considerable security against plant pathogen attack. However, despite \$1 billion worth of expenditure on fungicides, worldwide crop losses amounted to approximately 10% of crop value in 1981 (James, 1981; Seed Sci. & Technol. 9: 679-685).

The severity of the destructive process of disease depends on the aggressiveness of the pathogen and the response of the host. One aim of most plant breeding programs is to increase the resistance of host plants to disease. Typically, different races of pathogens interact with different varieties of the same crop species differentially, and many sources of host resistance only protect against specific pathogen races. Furthermore, some pathogen races show early signs of disease symptoms, but cause little damage to the crop. Jones and Clifford (1983; Cereal Diseases, John Wiley) report that virulent forms of the pathogen are expected to emerge in the pathogen population in response to the introduction of resistance into host cultivars and that it is therefore necessary to monitor pathogen populations. In addition, there are several documented cases of the evolution of fungal strains which are

resistant to particular fungicides. As early as 1981, Fletcher and Wolfe (1981; *Proc. 1981 Brit. Crop Prot. Conf.*) contended that 24% of the powdery mildew populations from spring barley, and 53% from winter barley showed considerable variation in response to the fungicide triadimenol and that the distribution of these populations varied between varieties with the most susceptible variety also giving the highest incidence of less susceptible types. Similar variation in the sensitivity of fungi to fungicides has been documented for wheat mildew (also to triadimenol), *Botrytis* (to benomyl), *Pyrenophora* (to organomercury), *Pseudocercosporella* (to MBC-type fungicides) and *Mycosphaerella fijiensis* to triazoles to mention just a few (Jones and Clifford; Cereal Diseases, John Wiley, 1983).

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Cereal species are grown world-wide and represent a major fraction of world food production. Although yield loss is caused by many pathogens, the necrotizing pathogens Septoria and Pseudocercosporella are particularly important in the major cereal growing areas of Europe and North America (Jones and Clifford; Cereal Diseases, John Wiley, 1983). In particular, the differential symptomology caused by different isolates and species of these fungi make the accurate predictive determination of potential disease loss difficult. Consequently, the availability of improved diagnostic techniques for the rapid and accurate identification of specific pathogens will be of considerable use to field pathologists.

Four Septoria species parasitize the small grain species. Septoria tritici is the causative agent 20 of leaf blotch and is virulent on wheat but also parasitizes triticale and rye. It typically causes leaf necrosis. Septoria nodorum is the causative agent of glume blotch and is parasitic on wheat, triticale, rye and barley and although mainly restricted to glumes is also found on leaf blades and sheaths. Septoria avenae is parasitic on oats, wheat and triticale and Septoria passerinii is restricted to barley. Septoria diseases occur in all wheat growing areas at 25 economically important levels. Different Septoria diseases frequently occur concurrently within fields and on individual plants, where the disease symptoms may be collectively referred to as the "Septoria complex". Typically, the most commonly found species are S. tritici and S. nodorum. According to Wiese (1977; Compendium of Wheat Diseases, Amer. Phytopath. Soc. pages 42-45), the Septoria complex presently destroys nearly 2% of the 30 world's wheat annually, the yield loss being mainly the result of impaired grain filling. Fungicide treatments can save up to 20% in cases of severe Septoria infection, but it is often

difficult to distinguish between the different Septoria species at the onset of infection and this

makes the decision whether or not to invest in fungicide use difficult because different cultivars display differing degrees of resistance to the various Septoria species.

The eyespot disease of cereals is caused by the fungus Pseudocercosporella herpotrichoides and is restricted to the basal culm of the plant. Wheat, rye, oats and other grasses are susceptible to the eyespot disease which occurs in cool, moist climates and is prevalent in Europe, North and South America, Africa and Australia. Wheat is the most susceptible cereal species, but isolates have been identified which are also virulent on other cereals. The R-strain of the fungus, for example, has also been isolated from rye and grows more slowly on wheat than the W-strain which has been isolated from wheat. Although eyespot may kill tillers or plants outright, it more usually causes lodging and/or results in a reduction in kernel size and number. Yield losses associated with eyespot are of even greater magnitude than those associated with Septoria tritici and Septoria nodorum. Typical control measures for eyespot include treatment with growth regulators to strengthen internodes, and fungicide treatment. However, the differing susceptibility of cultivars to different strains of the fungus render the predictive efficacy of fungicide treatments difficult.

Sigatoka leaf spot of banana occurs in two forms each of which is caused by a different fungus. The economically important Black Sigatoka is caused by Mycosphaerella fijiensis, whereas the less economically significant Yellow Sigatoka is caused by Mycosphaerella musicola (Johanson and Jeger, 1993; Mycol. Res. 97: 670-674). Black Sigatoka is the major problem in banana causing severe losses of 30% and more. Due to occurrence of fungicide resistance in Mycosphaerella fijiensis, usage of fungicide should best be limited to prevent the further occurrence of resistance. Consequently, the availability of diagnostic tools will provide an important means of identifying the appropriate circumstances in which to utilize fungicides without unnecessarily risking the development of further resistance.

Thus, there is a real need for the development of technology which will allow the identification of specific races of pathogen fungi early in the infection process. By identifying the specific race of a pathogen before disease symptoms become evident in the crop stand, the agriculturist can assess the likely effects of further development of the pathogen in the crop variety in which it has been identified and can choose an appropriate fungicide if such application is deemed necessary.

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#### **SUMMARY OF THE INVENTION**

The present invention is drawn to methods of identification of different pathotypes of plant pathogenic fungi. The invention provides DNA sequences which show variability between different fungal pathotypes. Such DNA sequences are useful in the method of the invention as they can be used to derive primers for use in polymerase chain reaction (PCR)-based diagnostic assays. These primers generate unique fragments in PCR reactions in which the DNA template is provided by specific fungal pathotypes and can thus be used to identify the presence or absence of specific pathotypes in host plant material before the onset of disease symptoms.

This invention provides the possibility of assessing potential damage in a specific crop variety-pathogen strain relationship and of utilizing judiciously the diverse armory of fungicides which is available. Furthermore, it can be used to provide detailed information on the development and spread of specific pathogen races over extended geographical areas. The invention provides a method of detection which is especially suitable for diseases with a long latent phase such as those caused by Septoria nodorum or Septoria tritici on wheat and Mycosphaerella fijiensis on banana.

Kits useful in the practice of the invention are also provided. The kits find particular use in the identification of Septoria, Pseudocercosporella, Fusarium, and Mycosphaerella pathogens.

#### **DESCRIPTION OF THE FIGURES**

Figure 1 Alignment of Internal Transcribed Spacer Sequences from Septoria tritici,

Septoria nodorum, Pseudocercosporella herpotrichoides strain W (two variants),

Pseudocercosporella herpotrichoides strain R, Mycosphaerella fijiensis, and Mycosphaerella
musicola.

Figure 2 Alignment of the Internal Transcribed Spacer Sequences from Septoria nodorum and Septoria avenae f.sp. triticea.

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Figure 3 Alignment of the Internal Transcribed Spacer Sequences from Fusarium graminearum, Fusarium culmorum, Fusarium moniliforme and Microdochium nivale.

### DETAILED DESCRIPTION OF THE INVENTION

The present invention provides unique DNA sequences which are useful in identifying different pathotypes of plant pathogenic fungi. Particularly the DNA sequences can be used as primers in PCR based analysis for the identification of fungal pathotypes. The DNA sequences of the invention include the Internal Transcribed Spacer (ITS) of the ribosomal RNA gene regions of particular fungal pathogens as well as primers which are derived from these regions which are capable of identifying the particular pathogen. These ITS DNA sequences from different pathotypes within a pathogen species or genus which vary between the different members of the species or genus can be used to identify those specific members.

Biomedical researchers have used PCR-based techniques for some time and with moderate success to detect pathogens in infected animal tissues. Only recently, however, has this technique been applied to detect plant pathogens. The presence of *Gaumannomyces graminis* in infected wheat has been detected using PCR of sequences specific to the pathogen mitochondrial genome (Schlesser et al., 1991; Applied and Environ. Microbiol. 57: 553-556) and random amplified polymorphic DNA (i.e. RAPD) markers were able to distinguish numerous races of Gremmeniella abietina, the causal agent of scleroderris canker in conifers.

Ribosomal genes are suitable for use as molecular probe targets because of their high copy number. Despite the high conservation between mature rRNA sequences, the non-transcribed and transcribed spacer sequences are usually poorly conserved and are thus suitable as target sequences for the detection of recent evolutionary divergence. Fungal rRNA genes are organized in units each of which encodes three mature subunits of 18S, 5.8S, and 28S respectively. These subunits are separated by two internal transcribed spacers, ITS1 and ITS2, of around 300 bp (White et al., 1990; In: PCR Protocols; Eds.: Innes et al.; pages 315-322). In addition, the transcriptional units are separated by non-transcribed spacer

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sequences (NTSs). The ITS and NTS sequences are particularly suitable for the detection of specific pathotypes of different fungal pathogens.

The DNA sequences of the invention are from the Internal Transcribed Spacer (ITS) of the ribosomal RNA gene region of different plant pathogens. The ITS DNA sequences from different pathotypes within a pathogen species or genus vary between the different members of the species or genus. Once having determined the ITS sequences of a pathogen, these sequences can be aligned with other ITS sequences. In this manner, primers can be derived from the ITS sequences. That is, primers can be designed based on regions within the ITS regions that contain the greatest differences in sequence among the fungal pathotypes. These sequences and primers based on these sequences can be used to identify specific pathogen members.

Particular DNA sequences of interest include ITS DNA sequences from Septoria, particularly, Septoria nodorum and Septoria tritici; Mycosphaerella, particularly Mycosphaerella fijiensis and Mycosphaerella musicola; Pseudocercosphorella, particularly Pseudocercosporella herpotrichoides, more particularly for the W-strain and the R-strain of Pseudocercosporella herpotrichoides, Fusarium, particularly F. graminearum, F. culmorum, F. moniliforme and Microdochium nivale. Such ITS DNA sequences as well as primers of interest are given in SEQ ID NO: 1 - 47 and SEQ ID NO.: 50-86. The sequences find use in 20 the PCR-based identification of the pathotypes of interest.

Methods for the use of the primer sequences of the invention in PCR analysis are well known in the art. For example, see US Patent Nos. 4,683,195 and 4,683,202 as well as Schlesser et al. (1991) Applied and Environ. Microbiol. 57:553-556. See also, Nazar et al. (1991; Physiol. and Molec. Plant Pathol. 39: 1-11) which used PCR amplification to exploit differences in the ITS regions of Verticillium albo-atrum and Verticillium dahliae and therefore distinguish between the two species; and Johanson and Jeger (1993; Mycol. Res. 97: 670-674) who used similar techniques to distinguish the banana pathogens Mycosphaerella fijiensis and Mycospharella musicola.

The ITS DNA sequences of the invention can be cloned from fungal pathogens by methods known in the art. In general, the methods for the isolation of DNA from fungal isolates are

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known. See, Raeder & Broda (1985) Letters in Applied Microbiology 2:17-20; Lee et al. (1990) Fungal Genetics Newsletter 35:23-24; and Lee and Taylor (1990) In: PCR Protocols: A Guide to Methods and Applications, Innes et al. (Eds.); pages 282-287.

Alternatively, the ITS regions of interest can be determined by PCR amplification. Primers to amplify the entire ITS region were designed according to White et al. (1990; In: PCR Protocols; Eds.: Innes et al. pages 315-322) and the amplified ITS sequence was subcloned into the pCRII cloning vector. The subcloned sequence included the lefthand ITS (ITS1), the righthand ITS (ITS2) as well as the centrally located 5.8S rRNA gene. This was undertaken for Septoria nodorum and Septoria tritici, numerous Pseudocercosporella isolates and Mycosphaerella fijiensis, Mycosphaerella musicola, Septoria avenae triticea, F. graminearum, F. culmorum, F. moniliforme and Microdochium nivale.

The ITS sequences were determined and within each pathogen group the sequences were compared to locate divergences which might be useful to test in PCR to distinguish the different species and/or strains. The sequences of the ITS regions which were determined are shown as Sequence ID's 1 to 6, 47, and 82-86 and also in Figures 1, 2 and 3. From the identification of divergences numerous primers were synthesized and tested in PCR-amplification. Templates used for PCR-amplification testing were firstly purified pathogen DNA, and subsequently DNA isolated from infected host plant tissue. Thus it was possible to identify pairs of primers which were diagnostic *i.e.* which identified one particular pathogen species or strain but not another species or strain of the same pathogen. Preferred primer combinations are able to distinguish between the different species or strains in infected host tissue *i.e.* host tissue which has previously been infected with a specific pathogen species or strain.

This invention provides numerous primer combinations which fulfill this criterion for different Septoria, Mycosphaerella, and Fusarium species and different strains of Pseudocercosporella. The primers of the invention are designed based on sequence differences among the fungal ITS regions. A minimum of one base pair difference between sequences can permit design of a discriminatory primer. Primers designed to a specific fungal DNA's ITS region can be used in combination with a primer made to a conserved sequence region within the ribosomal DNA's coding region to amplify species-specific PCR

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fragments. In general, primers should have a theoretical melting temperature between about 60 to about 70 degree C to achieve good sensitivity and should be void of significant secondary structure and 3' overlaps between primer combinations. Primers are generally at least about 5 to about 10 nucleotide bases.

The usefulness of cloned ITS sequences for the selection of primers for diagnostic purposes is largely due to their rapid evolutionary divergence. For example, W-type and R-type isolates of the pathogen *Pseudocercosporella herpotrichoides* were found to have divergent ITS sequences from which diagnostic primers were developed. However, the rapid divergence within the ITS sequence is apparent from the observation that two different sequence variants of the W-type were identified. The sequence identity within the W-type was 99.4 %, whereas that between W and R-types was 98.6 % suggesting a closer evolutionary relationship between the two W variants than was found between the W and the R-types. This closer relationship is also apparent from their similar host pathogenicity of the two isolates with divergent ITS sequences.

In addition to developing primers from ITS-derived sequences for PCR diagnosis of fungal isolates, the invention also encompasses the identification of primers from RAPD primer libraries which can distinguish between Septoria nodorum and Septoria tritici when used in PCR. The primers screened are commercially available and were obtained from Operon Technologies Incorporated (Alameda, CA). Screening on Septoria genomic DNA identified two primers which were able to detect only S. tritici and three which were able to detect only S. nodorum.

The present invention lends itself readily to the preparation of "kits" containing the elements necessary to carry out the process. Such a kit may comprise a carrier being compartmentalized to receive in close confinement therein one or more container means, such as tubes or vials. One of said container means may contain unlabeled or detectably labeled DNA primers. The labeled DNA primers may be present in lyophilized form, or in an appropriate buffer as necessary. One or more container means may contain one or more enzymes or reagents to be utilized in PCR reactions. These enzymes may be present by themselves or in admixtures, in lyophilized form or in appropriate buffers.

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Finally, the kit may contain all of the additional elements necessary to carry out the technique of the invention, such as buffers, extraction reagents, enzymes, pipettes, plates, nucleic acids, nucleoside triphosphates, filter paper, gel materials, transfer materials, autoradiography supplies, and the like.

The examples below show, without limitation, typical experimental protocols which can be used in the isolation of ITS sequences, the selection of suitable primer sequences, the testing of primers for selective and diagnostic efficacy, and the use of such primers for disease and fungal isolate detection. Such examples are provided by way of illustration and not by way of limitation.

#### **EXAMPLES**

# 15 Example 1: Fungal isolates and genomic DNA extraction

Viable fungal isolates of S. nodorum, S. tritici, S. passerini, S. glycines, Pseudocercosporella herpotrichoides, Pseudocercosporella aestiva, Mycosphaerella citri, Mycosphaerella graminicola, Mycosphaerella fijiensis and Mycosphaerella musicola were obtained from the American Type Culture Collection. Fusarium culmorum and Fusarium graminearum isolates were obtained from Dr. Paul Nelson from Penn State University. An isolate of Michrodochium nivale (syn. Fusarium nivale) was received from Ciba- Basel and an isolate of Fusarium moniliforme was received from Dr. Loral Castor. Fungi were grown in 150 ml potato dextrose broth inoculated with mycelial fragments from PDA (Potato Dextrose Agar) cultures. Cultures were incubated on an orbital shaker at 28°C for 7-11 days. Mycelia were pelleted by centrifugation and then ground in liquid nitrogen and total genomic DNA extracted using the protocol of Lee and Taylor (1990; In: PCR Protocols: A Guide to Methods and Applications; Eds.: Innes et al.; pages 282-287).

Dr. Bruce McDonald from Texas A&M University supplied genomic DNA from ten isolates of S. nodorum and nine isolates of S. tritici. Dr. Chris Caten of Birmingham University provided six isolates of Septoria nodorum purified fungal DNA. Purified genomic DNA from 12 isolates of Pseudocercosporella herpotrichoides was obtained from Dr. Paul Nicholson of the John Innes Centre, Norwich, UK. Six of these isolates are of the W-type;

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the other six isolates are of the R-type. These isolates were typed based on pathogenicity and RFLP studies. Andrea Johanson of the Natural Resources Institute supplied genomic DNA of six isolates of *M. musicola*, six isolates of *M. fijiensis* and a single isolate of *Mycosphaerella musae*. Purified genomic DNA from *Septoria avenae* f. sp. triticea ATCC#26380 was supplied by Dr. Peter Ueng from the USDA at Beltsville, Maryland.

Table 1: Source of Test Isolates

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<u>Isolate</u>	<u>Species</u>	<u>Origin</u>	Source
ATCC#24425	S. nodorum	Montana	ATCC1
XA1.1	S. nodorum	Texas	B. McDonald
Xa5A.2	S. nodorum	Texas	B. McDonald
YA3.1	S. nodorum	Texas	B. McDonald
XD2.1	S. nodorum	Texas	B. McDonald
YB2.2	S. nodorum	Texas	B. McDonald
93HBh6a	S. nodorum	Oregon	B. McDonald
93A3a	S. nodorum	Oregon	B. McDonald
93AYa	S. nodorum	Oregon	B. McDonald
93HBh8a	S. nodorum	Oregon	B. McDonald
93C5a	S. nodorum	Oregon	B. McDonald
ATCC#26517	S. tritici	Minnesota	ATCC
BS3	S. nodorum	Ireland	C. Caten <sup>3</sup>
BS6	S. nodorum	Ireland	C. Caten
BS175	S. nodorum	<b>England</b>	C. Caten
BS425	S. nodorum	England	C. Caten
alpha'5	S. nodorum	France	C. Caten
m300	S. nodorum	England	C. Caten
TKV2a	S. tritici	Turkey	B. McDonald
SYK2	S. tritici	Syria	B. McDonald
ISZC36.2	S. tritici	Israel	B. McDonald
CNRC4a.1	S. tritici	Canada	B. McDonale
ALA1a	S. tritici	Algeria	B. McDonald
ETK1	S. tritici	Ethiopia	B. McDonale
GEB2a.1	S. tritici	Germany	B. McDonald
UK92D2	S. tritici	United Kingdom	B. McDonale
DNB1a	S. tritici	Denmark	B. McDonale
ATCC#38699	S. glycines	Illinois	ATCC
ATCC#22585	S. passerini	Minnesota	ATCC
ATCC#42040	P. herpotrichoides-wheat		ATCC
ATCC#62012	P. aestiva	Germany	ATCC '
ATCC#60972	P. herp. var. herp barley	Germany	ATCC
W1	P. herpotrichoides	United Kingdom	P. Nicholson
W2	P. herpotrichoides	United Kingdom	P. Nicholson
W3	P. herpotrichoides	United Kingdom	P. Nicholson
W4	P. herpotrichoides	United Kingdom	P. Nicholson

<b>W</b> 5	P. herpotrichoides	New Zealand	P. Nicholson
W6	P. herpotrichoides	Italy	P. Nicholson
R1	P. herpotrichoides	Belgium	P. Nicholson
R2	P. herpotrichoides	New Zealand	P. Nicholson
R3	P. herpotrichoides	Germany	P. Nicholson
R4	P. herpotrichoides	Sweden	P. Nicholson
R5	P. herpotrichoides	United Kingdom	P. Nicholson
R6	P. herpotrichoides	United Kingdom	P. Nicholson
ATCC#22116	M. fijiensis	Philippines	ATCC
ATCC#22115	M. musicola	Philippines	ATCC
ATCC#24046	M. citri	Florida	ATCC
ATCC#62714	M. graminicola	Montana	ATCC
PA92	M. fijiensis	Panama	A. Johanson <sup>5</sup>
PNG291	M. fijiensis	Papua New Guinea	A. Johanson
GH6-3	M. fijiensis	Ghana	A. Johanson
TG120	M. fijiensis	Tonga	A. Johanson
HSB4	M. fijiensis	Honduras	A. Johanson
RT689	M. fijiensis	Rarotonga (Cook Is.)	A. Johanson
CR548	M. musicola	Costa Rica	A. Johanson
CM61	M. musicola	Cameroon	A. Johanson
CU823	M. musicola	Cuba	A. Johanson
MQ103	M. musicola	Martinique	A. Johanson
CI31	M. musicola	Ivory Coast	A. Johanson
CB90	M. musicola	Colombia	A. Johanson
BD1-4	M. musae	Barbados	A. Johanson
ATCC#44234	Ceratobasidium cereale	Netherlands	ATCC
ATCC#11404	Drechslera sorokiniana	Minnesota	ATCC
R-5126	F. culmorum	Minnesota	P. Nelson <sup>0</sup>
R-5106	F. culmorum	Michigan	P. Nelson
R-5146	F. culmorum	Finland	P. Nelson
R-8417	F. graminearum	Italy	P. Nelson
R-8422	F. graminearum	Canada	P. Nelson
R-8546	F. graminearum	Bulgaria	P. Nelson
4551	F. moniliforme	Indiana	L. Castor
92	M. nivale	******	Ciba Başel
ATCC#26380	S. avenae f.sp.triticea	Minnesota	P. Ueng

<sup>1</sup> American Type Culture Collection, Rockville, Maryland USA

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<sup>2</sup> Dr. Bruce McDonald, Texas A&M University, USA

<sup>3</sup> Dr. Chris Caten, Birmingham University, UK

<sup>4</sup> Dr. Paul Nicholson, John Innes Centre, UK

<sup>5</sup> Dr. Andrea Johanson, Natural Resources Institute, UK

<sup>6</sup> Dr. Paul Nelson, Penn State University

<sup>7</sup> Dr. Loral Castor, Ciba Seeds Research, Bloomington, Illinois

<sup>8</sup> Ciba-Geigy Limited, Basel, Switzerland

<sup>9</sup> Dr. Peter Ueng, USDA, Beltsville, Maryland

Example 2: Isolation of the internal transcribed spacer (ITS) regions

The approximately 550 bp internal transcribed spacer region fragments were PCR amplified from 25 ng of genomic DNA isolated from S. nodorum (ATCC#24425), S. tritici (ATCC#26517), Pseudocercosporella herpotrichoides isolates R1, R2, W2 and W5, M.

fijiensis (ATCC#22115) and M. musicola (ATCC#22115) using 50 pmol of primers ITS1 (5'-TCCGTAGGTGAACCTGCGG-3'; SEQ ID NO: 38) and ITS4 (5'-TCCTCCGCTTATTGATATGC-3'; SEQ ID NO:41). PCRs were performed as described in EXAMPLE 4 except that reactions were done in 100 μl and annealing was done at of 50°C. The ITS fragments were purified by isopropanol precipitation according to Maniatis et al.

in 50 µl dH<sub>2</sub>O and cloned using the Invitrogen Corporation's (San Diego, CA) TA Cloning Kit (part no. K2000-01) using the pCRII cloning vector. The DNA sequences of the ITS regions were determined by the dideoxy method using the Applied Biosystems (Foster City, CA) automated sequencer model 373A with the primers ITS1 (see sequence above), ITS2 (5'-GCTGCGTTCTTCATCGATGC-3'; SEQ ID NO:39), ITS4 (see sequence above) and the M13 universal -20 (5'-GTAAAACGACGGCCAGT-3'; SEQ ID NO:48) and Reverse (5'-AACAGCTATGACCATG-3'; SEQ ID NO:49) primers. The ITS primers ITS1 (SEQ ID NO:38), ITS2 (SEQ ID NO:39), ITS3 (SEQ ID NO:40), and ITS4 (SEQ ID NO:41) used for cloning the ITS regions are detailed in White *et al.* (1990; In: PCR Protocols; Eds.: Innes *et* 

(1982; Molecular Cloning; Eds.: Maniatis et al.; pages 461-462). The DNA was resuspended

In addition, the internal transcribed spacer regions were PCR amplified from 25 ng of genomic DNA from S. avenae f.sp. triticea, M. nivale, F. moniliforme (#4551), F. graminearum isolates R-8417, R-8546 and R-8422 and F. culmorum isolates R-5126, R-5106 and R-5146. PCR products were purified using Promega's Wizard DNA Clean-up kit (Madison, WI). The DNA sequences of the ITS regions were determined as described above using the ITS1 (SEQ ID NO:38), ITS2 (SEQ ID NO:39), ITS3 (SEQ ID NO:40) and ITS4 (SEQ ID NO:41) primers. Sequencing reactions were combined with the three isolates of F. culmorum and F. graminearum to generate a consensus sequence for F. culmorum and F. graminearum.

## Example 3: DNA extraction from wheat and banana leaves

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al. pages 315-322).

DNA was extracted from wheat leaves using a modified version of the Rapid DNA Extraction protocol from the MicroProbe Corporation's (Garden Grove, CA) IsoQuick Nucleic Acid Extraction Kit (cat# MXT-020-100). Typical yields were 5-10 µg of total DNA from 0.2 g of leaf tissue. Approximately 100 ng of total DNA were used in each PCR assay.

### Modified Rapid DNA Extraction:

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Before using kit for the first time, the entire contents of Reagent 2A (20 x Dye Concentrate)
were added to Reagent 2 (Extraction Matrix).

- (1) Approximately 0.2 g of leaf sample were added to a 1.5 ml eppendorf tube containing 50  $\mu$ l sample buffer A and 50  $\mu$ l #1 lysis solution. The leaf sample was ground with a Kontes pestle.
- 15 (2) Reagent 2 (Extraction Matrix) was shaken vigorously. 350 μl of reagent 2 were added to the sample lysate.
  - (3) 200  $\mu$ l of Reagent 3 were added (Extraction Buffer) to the sample. The sample was vortexed 20 sec.
  - (4) Microcentrifugation at 12,000 x g for 5 min.
- 20 (5) The aqueous phase (upper layer) was transferred to a new microcentrifuge tube. This volume was typically about 200  $\mu$ l.
  - (6) 0.1 x the volume of the aqueous phase of Reagent 4 (Sodium Acetate) to the aqueous phase sample.
- (7) An equal volume of isopropanol was added to the aqueous phase sample followed by vortexing.
  - (8) Microcentrifugation at 12,000 x g for 10 min.
  - (9) The supernatant was discarded without disturbing the nucleic acid pellet. 0.5 ml of -20°C 70% ethanol was added to the pellet. The tube was vortexed to mix.
  - (10) Microcentrifugation at 12,000 x g for 5 min.
- 30 (11) The supernatant was discarded and the pellet was allowed to dry.
  - (12) The nucleic acid pellet was dissolved in 50  $\mu$ l Reagent 5 (RNase-free water).

# Example 4: Polymerase chain reaction amplification

Polymerase chain reactions were performed with the GeneAmp Kit from Perkin-Elmer/Cetus (Norwalk, CT; part no. N808-0009) using 50 mM KCl, 2.5 mM MgCl<sub>2</sub>, 10 mM Tris-HCl, pH8.3, containing 100 μM of each TTP, dATP, dCTP, and dGTP, 50 pM primer, 2.5 units of Taq polymerase and 25 ng of genomic DNA in a final volume of 50 μl. Reactions were run for 30 cycles of 15 s at 94°C, 15 s at 50°C, 60°C or 70°C, and 45 s at 72°C in a Perkin-Elmer/Cetus Model 9600 thermal cycler. The products were analyzed by loading 20 μl of each PCR sample on a 1.1-1.2% agarose gel and electrophoresed.

# 10 Example 5: Synthesis and Purification of Oligonucleotides

Oligonucleotides (primers) were synthesized on an Applied Biosystems 380A DNA synthesizer using B-cyanothyl-phosphoramidite chemistry.

# 15 Example 6: Selection of species-specific primers

The ITS sequences of S. nodorum, S. tritici, P. herpotrichoides strains R and W, M. fijiensis and M. musicola were aligned (Fig. 1). The ITS sequences of S. nodorum and S. avenae. triticea were aligned (Fig. 2). An alignment was also made of the ITS sequences from F. graminearum, F. culmorum, F. moniliforme and M. nivale (Fig. 3). Sets of primers were synthesized according to EXAMPLE 5 based on analysis of the aligned sequences. Primers were designed to regions containing the greatest differences in sequence among the fungal species for Figs. 1-2. In Fig 3, primers were designed to regions of highest homology within the ITS for Fusarium. In addition, the published ribosomal gene-specific primers ITS1 (SEQ ID NO:38), ITS2 (SEQ ID NO:39), ITS3 (SEQ ID NO:40) and ITS4 (SEQ ID NO:41) (White et al., 1990; In: PCR Protocols; Eds.: Innes et al. pages 315-322) were synthesized for testing in combination with the primers specific for the ITS region.

Table 2: Primer Design for Fungal Detection

)	Primer Template	Primer	Name Primer Sequence
	S. nodorum	ЈВ433	5' ACACTCAGTAGTTTACTACT 3' (SEQ ID NO:7)
	S. nodorum	ЈВ434	5' TGTGCTGCGCTTCAATA 3' (SEQ ID NO:8)
	S. nodorum	ЈВ525	5' GCGACTTGTGCTGCGCTTCAATA 3' (SEQ ID NO:9)

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5' CATTACACTCAGTAGTTTACTACT 3' (SEQ ID NO:10)
               JB527
S. nodorum
                      5' CTGCGTCGGAGTTTACG 3' (SEQ ID NO:11)
               JB445
S. tritici
                      5' CGAGGCTGGAGTGGTGT 3' (SEQ ID NO:12)
               JB446
S. tritici
                      5' CCCAGCGAGGCTGGAGTGGTGT 3' (SEQ ID NO:13)
               JB526
S. tritici
                      5' CTGGGGGCTACCTACTTGGTAG 3' (SEQ ID NO:14)
               JB536
P. herp.
                      5' GGGGGCTACCCTACTTGGTAG 3' (SEQ ID NO:15)
               JB537
P. herp.
                      5' ACTTGGTAGGGTTTAGAGTCGTCA 3' (SEQ ID NO:16)
               JB538
P. herp.
                      5' CTTCGGTAAGGTTTAGAGTCGTCG 3' (SEQ ID NO:17)
               JB539
P. herp.
                      5' GGGGGCCACCCTACTTCGGTAA 3' (SEQ ID NO:18)
               JB540
P. herp.
                      5' CCACTGATTTTAGAGGCCGCGAG 3' (SEQ ID NO:19)
               JB541
P. herp.
                      5' CCACTGATTTTAGAGGCCGCGAA 3' (SEQ ID NO:20)
P. herp.
               JB542
                      5' CCTGTAAAAATTGGGGGTTA 3' (SEQ ID NO:21)
               JB543
P. herp.
                       5' CCTGTAAAAAATTGGGGGTTG 3' (SEQ ID NO:22)
               JB544
P. herp.
                       5' ATTACCGAGTGAGGGCTCACGC 3' (SEQ ID NO:23)
               JB547
M. fijiensis
                       5' GTTGCTTCGGGGGCGACCTG 3' (SEQ ID NO:24)
               JB548
M. fijiensis
                       5' TCGGGGGCGACCTGCCG 3' (SEQ ID NO:25)
               JB442
M. fijiensis
                       5' CCGGAGGCCGTCTA 3' (SEQ ID NO:26)
               JB443
M. fijiensis
                       5' CCACAACGCTTAGAGACGGACAG 3' (SEQ ID NO:27)
               JB545
M. fijiensis
                       5' CACCCGCACTCCGAAGCGAATT 3' (SEQ ID NO:28)
               JB546
M. fijiensis
                       5' GATCCGAGGTCAACCTTTGAATAA 3' (SEQ ID NO:29)
               JB549
M. fijiensis
                       5' GGTCAACCTTTGAATAA 3' (SEQ ID NO:30)
               JB444
M. fijiensis
                       5' CCTTTGTGAACCACACCT 3' (SEQ ID NO:31)
               JB451
M. musicola
                       5' CTGCCGGCGAACTT 3' (SEQ ID NO:32)
               JB440
M. musicola
                       5' ACCCTGCCGGCGAACTT 3' (SEQ ID NO:33)
               JB449
M. musicola
                       5' GCGACCCTGCCGGCGAAC 3' (SEQ ID NO:34)
               JB448
M. musicola
                       5' TAGCCGGGAGACTTTGG 3' (SEQ ID NO:35)
M. musicola.
               JB441
                       5' TCTGCGTCGGAGTTCC 3' (SEQ ID NO:36)
               JB450
M. musicola
                       5' CCGCGCTCCGGAGCGAAC 3' (SEQ ID NO:37)
               JB452
M. musicola
                       5' TCCGTAGGTGAACCTGCGG 3' (SEQ ID NO:38)
18S rDNA
               ITS1
                       5' GCTGCGTTCTTCATCGATGC 3' (SEQ ID NO:39)
 5.8S rDNA
               ITS2
                       5' GCATCGATGAAGAACGCAGC 3' (SEQ ID NO:40)
                ITS3
 5.8S rDNA
                       5' TCCTCCGCTTATTGATATGC 3' (SEQ ID NO:41)
                ITS4
 25S rDNA
                       5' CTTGCCTGCCGGTTGGACAAATT 3' (SEQ ID NO:50)
 S. nodorum
                JB563
                       5' CTCAGTAGTTTACTACTGTAAAAGG 3' (SEQ ID NO:51)
                JB564
 S. nodorum
                       5' CTTCTGGACGCAAGTGTTTGTTAC 3' (SEQ ID NO:52)
                JB565
 S. nodorum
                       5' GTTTTTAGTGGAACTTCTGAGT 3' (SEQ ID NO:53)
                JB566
 Fusarium spp.
                       5' CGCAGGAACCCTAAACTCT 3' (SEQ ID NO:54)
                JB567
 Fusarium spp.
                       5' GCCCGCCGCAGG 3' (SEQ ID NO:55)
                JB568
 Fusarium spp.
                       5' RTWWTTWRTGGAMYYTCTGAGT 3' (SEQ ID NO:56)
                JB569
 Fusarium spp.
                        5' TATGTTGCCTCGGCGG 3' (SEQ ID NO:57)
                JB570
 Fusarium spp.
                       5' TAACGATATGTAAATTACTACGCT 3' (SEQ ID NO:58)
                JB571
 Fusarium spp.
                       5' AAGTTGGGGTTTAACGGC 3' (SEQ ID NO:59)
                JB572
 Fusarium spp.
                       5' AGCGAGCCCGCCAC 3' (SEQ ID NO:60)
                JB573
 Fusarium spp.
                        5' CCATTGTGAACGTTACCTATAC 3' (SEQ ID NO:61)
 Fusarium spp.
                JB574
                        5' CGACCAGAGCGAGATGTA 3' (SEQ ID NO:62)
                JB575
 Fusarium spp.
                        5' GTGAACATACCTTATGTTGCC 3' (SEQ ID NO:63)
                JB576
 Fusarium spp.
                        5' GTTGCCTCGGCGGATC 3' (SEQ ID NO:64)
                JB577
 Fusarium spp.
                        5' CCGCGACGATTACCAG 3' (SEQ ID NO:65)
                JB578
 Fusarium spp.
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NOTE: Fusarium spp. includes F. graminearum, F. culmorum, F. moniliforme and Michrodochium nivale (syn. F. nivale).

# Example 7: Selection of Random Amplified Polymorphic DNA (RAPD) primers

Two RAPD primer libraries (kits B and E) of twenty oligonucleotides each were purchased from Operon Technologies Incorporated (Alameda, CA). The primers were tested for their ability to differentiate purified genomic DNA of S. nodorum, S. tritici, M. fijiensis and M. musicola. The PCR conditions were essentially the same as described in EXAMPLE 4 except the number of PCR cycles was increased to 35, the annealing temperature was 30°C and only 5 picamoles of each primer were used. Five RAPD primers were identified that differentiate purified genomic DNA of S. nodorum, S. tritici, M. fijiensis and M. musicola. Primers OPB-12 and OPE-6 produced a single fragment when amplified with S. tritici genomic DNA. Primers OPB-12, OPB-19 and OPE-15 produced single fragments from S. nodorum genomic DNA. Primers OPB-12 and OPE-6 did not produce any amplification products from S. nodorum M. fijiensis and M. musicola genomic DNA. Primers OPE-12, OPB-19 and OPE-15 did not amplify any fragments from genomic S. tritici, M. fijiensis or M. musicola DNA.

Table 3: RAPD Primers for Septoria Diagnosis

25	Source of template DNA	<u>Primer</u>	Sequence of primer	Approximate size of amplified fragment
30	S. tritici S. tritici S. nodorum S. nodorum S. nodorum	OPB-12 OPE-6 OPE-12 OPB-19 OPE-15	5'-CCTTGACGCA-3' (SEQ ID NO: 42) 5'-AAGACCCCTC-3' (SEQ ID NO: 43) 5'-TTATCGCCCC-3' (SEQ ID NO: 44) 5'-ACCCCCGAAG-3' (SEQ ID NO: 45) 5'-ACGCACAACC-3' (SEQ ID NO: 46)	) 1.0 kb 2.2 kb ) 1.1 kb

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# Example 8: Determination of primer specificity to purified fungal genomic DNA

PCRs were performed according to EXAMPLE 4 using different primer combinations in an attempt to amplify a single species-specific fragment. Species-specific PCR amplification products were produced from primers designed from the ITS region between the 18S and 25S ribosomal DNA subunits of each fungal strain of interest.

Table 4: ITS-derived diagnostic PCR primers

S		Ар	proximate size of amplified
Source of template DNA	<u>5'Primer</u>	3'Primer	fragment
Septoria nodorum	JB433 (SEQ ID NO:7)	JB434 (SEQ ID NO:8)	448bp
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	JB433 (SEQ ID NO:7)	ITS4 (SEQ ID NO:41)(JB415)	553bp
	ITS1 (SEQ ID NO:38)(JB410)	JB434 (SEQ ID NO:8)	478bp
	ITS3 (SEQ ID NO:40)(JB414)	JB434 (SEQ ID NO:8)	232bp*
	JB527 (SEQ ID NO:10)	JB525 (SEQ ID NO:9)	458bp
	JB564 (SEQ ID NO:51)	JB565 (SEQ ID NO:52)	480bp
	JB563 (SEQ ID NO:50)	JB565 (SEQ ID NO:52)	368bp
	•		
Septoria tritic			4071
_	JB445 (SEQ ID NO:11)	ITS4 (SEQ ID NO:41)(JB415)	407bp
	ITS1 (SEQ ID NO:38)(JB410)	JB446 (SEQ ID NO:12)	345bp
	ITS3 (SEQ ID NO:40)(JB414)	JB446 (SEQ ID NO:12)	143bp*
	JB445 (SEQ ID NO:11)	JB446 (SEQ ID NO:12)	204bp
*			•
M. fijiensis			41.01
	JB443 (SEQ ID NO:26)	ITS4 (SEQ ID NO:41)(JB415)	418bp
	ITS1 (SEQ ID NO:38)(JB410)	JB444 (SEQ ID NO:30)	482bp
	JB443 (SEQ ID NO:26)	JB444 (SEQ ID NO:30)	366bp*
	ITS3 (SEQ ID NO:40)(JB414)	JB444 (SEQ ID NO:30)	281bp*
	ITS1 (SEQ ID NO:38)(JB410)	JB549 (SEQ ID NO:29)	489bp
			•
M. musicola		•	
M. musicoia			
	JB449 (SEQ ID NO:33)	ITS4 (SEQ ID NO:41)(JB415)	
	JB448 (SEQ ID NO:34)	ITS4 (SEQ ID NO:41)(JB415)	449bp*
	JB448 (SEQ ID NO:34)	ITS2 (SEQ ID NO:39)(JB411)	138bp*
	JB450 (SEQ ID NO:36)	ITS4 (SEQ ID NO:41)(JB415)	390bp*

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P. herpotrichoides JB536 (SEQ ID NO:14)	JB541 (SEQ ID NO:19)	415bp <sup>+</sup>
JB536 (SEQ ID NO:14)	JB543 (SEQ ID NO:21)	502bp
JB537 (SEQ ID NO:15)	JB541 (SEQ ID NO:19)	413bp <sup>+</sup>
JB537 (SEQ ID NO:15)	JB543 (SEQ ID NO:21)	500bp <sup>+</sup>
JB538 (SEQ ID NO:16)	JB541 (SEQ ID NO:19)	401bp
JB538 (SEQ ID NO:16)	JB543 (SEQ ID NO:21)	488bp
JB536 (SEQ ID NO:14)	ITS4 (SEQ ID NO:41)(JB415)	560bp
JB530 (SEQ ID NO:14)  JB537 (SEQ ID NO:15)	ITS4 (SEQ ID NO:41)(JB415)	558bp <sup>+</sup>
JB538 (SEQ ID NO:16)	ITS4 (SEQ ID NO:41)(JB415)	546bp
ITS1 (SEQ ID NO:38)(JB410)	JB541 (SEQ ID NO:19)	482bp <sup>+</sup>
ITS1 (SEQ ID NO:38)(JB410)	JB543 (SEQ ID NO:21)	569bp <sup>+</sup>
ITS1 (SEQ ID NO:38)(JB410)	JB542 (SEQ ID NO:20)	482bp <sup>+</sup>
ITS1 (SEQ ID NO:38)(JB410)	JB544 (SEQ ID NO:22)	569bp <sup>↔</sup>
JB540 (SEQ ID NO:18)	ITS4 (SEQ ID NO:41)(JB415)	. 558bp <sup>↔</sup>
JB539 (SEQ ID NO:17)	ITS4 (SEQ ID NO:41)(JB415)	545bp <sup>↔</sup>
JB540 (SEQ ID NO:18)	JB542 (SEQ ID NO:20)	413bp <sup>™</sup>
JB540 (SEQ ID NO:18)	JB544 (SEQ ID NO:22)	500bp**
JB539 (SEQ ID NO:17)	JB542 (SEQ ID NO:20)	400bp**
JB539 (SEQ ID NO:17)	JB544 (SEQ ID NO:22)	487bp <sup>™</sup>
	•	
Fusarium spp.		
JB566 (SEQ ID NO:53)	ITS4 (SEQ ID NO:41)(JB415)	430bp'
JB566 (SEQ ID NO:53)	JB572 (SEQ ID NO:59)	346bp <sup>1</sup>
JB569 (SEQ ID NO:56)	ITS4 (SEQ ID NO:41)(JB415)	430bp
JB569 (SEQ ID NO:56)	JB572 (SEQ ID NO:59)	346bp <sup>1</sup>
ITS1 (SEQ ID NO:38)(JB410)	JB572 (SEQ ID NO:59)	485bp¹
JB566 (SEQ ID NO:53)	JB571 (SEQ ID NO:58)	308bp²
JB569 (SEQ ID NO:56)	JB571 (SEQ ID NO:58)	308bp²
JB570 (SEQ ID NO:57)	ITS4 (SEQ ID NO:41)(JB415)	501bp <sup>2</sup>
JB570 (SEQ ID NO:57)	JB571 (SEQ ID NO:58)	379bp <sup>2</sup>
JB570 (SEQ ID NO:57)	JB578 (SEQ ID NO:65)	395bp <sup>2</sup>
JB567 (SEQ ID NO:54)	ITS4 (SEQ ID NO:41)(JB415)	450bp <sup>2</sup>
JB567 (SEQ ID NO:54)	JB571 (SEQ ID NO:58)	328bp <sup>2</sup>
JB567 (SEQ ID NO:54)	JB572 (SEQ ID NO:59)	366bp <sup>2</sup>
JB567 (SEQ ID NO:54)	JB578 (SEQ ID NO:65)	344bp <sup>2</sup>
JB568 (SEQ ID NO:55)	ITS4 (SEQ ID NO:41)(JB415)	459bp <sup>2</sup>
JB568 (SEQ ID NO:55)	JB571 (SEQ ID NO:58)	337bp <sup>2</sup>
JB568 (SEQ ID NO:55)	JB572 (SEQ ID NO:59)	375bp <sup>2</sup>
JB576 (SEQ ID NO:63)	ITS4 (SEQ ID NO:41)(JB415)	510bp <sup>2</sup>
JB576 (SEQ ID NO:63)	JB578 (SEQ ID NO:65)	404bp <sup>2</sup>
JB577 (SEQ ID NO:64)	ITS4 (SEQ ID NO:41)(JB415)	495bp <sup>2</sup>
JB577 (SEQ ID NO:64)	JB571 (SEQ ID NO:58)	373bp <sup>2</sup>
JB577 (SEQ ID NO:64)	JB578 (SEQ ID NO:65)	389bp <sup>2</sup>
ITS1 (SEQ ID NO:38)(JB410)	JB571 (SEQ ID NO:58)	447bp <sup>2</sup>
ITS1 (SEQ ID NO:38)(JB410)	JB578 (SEQ ID NO:65)	463bp <sup>2</sup>
ITS1 (SEQ ID NO:38)(JB410)	JB575 (SEQ ID NO:62)	479bp <sup>2</sup>
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<sup>\*...</sup> Primer combination amplified some fragments by false priming but none were the size of the desired fragment.

\*.... Primers amplified the correct size fragment from both R-type and W-type of Pseudocercosporella herpotrichoides.

\*\*... Primer combination amplified the correct size fragment from the R-type of P. herpotrichoides only.

<sup>1</sup>.... Primer combination amplified the correct size fragment from F. graminearum, F. culmorum, F. moniliforme and M. nivale.

<sup>2</sup>····· Primer combination amplified the correct size fragment from F. graminearum, F. culmorum and F. moniliforme.

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Example 9: Determination of primer specificity to plant tissue infected with fungi
Total genomic DNA was isolated from healthy wheat leaves, wheat leaves infected with S.

nodorum, wheat leaves infected with S. tritici and wheat leaves infected with both S.

nodorum and S. tritici using the protocol described in EXAMPLE 3. PCRs were performed as described in EXAMPLE 4 testing the primer combinations listed in EXAMPLE 8 against DNA from the wheat leaves.

The S. tritici-specific primer JB446 (SEQ ID NO:12) and ITS1 (SEQ ID NO:38)(JB410) amplified a 345 bp fragment from purified S. tritici DNA, from S. tritici-infected wheat leaf tissue and from a wheat leaf sample infected with both S. tritici and S. nodorum. The primer set did not amplify a diagnostic fragment from healthy wheat leaf tissue nor from S. nodorum-infected wheat tissue. Similarly, the S. tritici-specific primers JB445 (SEQ ID NO:11) and ITS4 (SEQ ID NO:41)(JB415) amplified a 407 bp fragment from the same tissues as the primer combination JB446 (SEQ ID NO:12) and ITS1 (SEQ ID NO:38)(JB410) and was also diagnostic.

Similarly diagnostic results were obtained with the S. nodorum-specific primers JB433 (SEQ ID NO:7) and JB434 (SEQ ID NO:8). The primers amplified a 448 bp fragment from S. nodorum-infected wheat tissue, from a wheat leaf sample infested with both S. nodorum and S. tritici, as well as from purified genomic DNA of S. nodorum. The primer combination

JB433 (SEQ ID NO:7) and JB434 (SEQ ID NO:8) did not amplify any fragments from healthy wheat tissue, from S.tritici-infected wheat tissue or from purified genomic DNA of S. tritici. The S. nodorum-specific primers JB527 (SEQ ID NO:10) and JB525 (SEQ ID NO:9) amplified a 458 bp fragment from the same genomic DNAs and wheat tissues as the JB433 (SEQ ID NO:7) and JB434 (SEQ ID NO:8) combination.

The P. herpotrichoides primer combinations listed in EXAMPLE 8 were PCR tested against the extracts from wheat stems as pbtained in Example 12. PCRs were performed as described in EXAMPLE 4 with the following changes: 35 cycles were run of 94°C for 15 sec and 70°C for 45 sec, 1.5 - 2.5 mM MgCl2 and 200 µM of each dNTP was used. 1 µl of wheat extract was used in each PCR.

Primer combination JB537 (SEQ ID NO:15) and JB541 (SEQ ID NO:19) amplified a 413 bp fragment from wheat extract infected with the W-type pathotype of *P. herpotrichoides*. No amplification products were produced from amplification with healthy wheat extract nor from wheat extract infected with the R-type pathotype of *P. herpotrichoides*.

The primer combination JB539 (SEQ ID NO:17) and JB544 (SEQ ID NO:22) amplified a 487 bp fragment and primer combination JB540 (SEQ ID NO:18) and JB542 (SEQ ID NO:20) amplified a 413 bp fragment from R-type infected wheat but not from healthy wheat nor from W-type infected wheat.

Total genomic DNA was also isolated from healthy banana leaves and from banana leaves infected with *M. fijiensis* using the protocol described in EXAMPLE 3. PCRs were performed as described in EXAMPLE 4 testing the *M. fijiensis* primer combinations listed in EXAMPLE 8 against DNA from the banana leaves.

The M. fijiensis-specific primer JB549 (SEQ ID NO:29) and ITS1 (SEQ ID NO:38)(JB410) amplified a 489 bp fragment from purified M. fijiensis DNA and from M. fijiensis-infected banana leaf tissue. The primer set did not amplify a diagnostic fragment from healthy banana leaf tissue. The M. fijiensis-specific primer combinations JB443 (SEQ ID NO:26)/ITS4 (SEQ ID NO:41)(JB415) and ITS1 (SEQ ID NO:38)(JB410)/JB444 (SEQ ID NO:30) amplified a 418 bp fragment and a 482 bp fragment, respectively, from the same genomic

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DNA and banana leaf tissue as the JB549 (SEQ ID NO:29) and ITS1 (SEQ ID NO:38)(JB410) primer combination.

5 Example 10: Determination of cross-reactivity of species-specific primers with other species and isolates

Purified fungal genomic DNAs were obtained as described in EXAMPLE 1 and PCR assayed as described in EXAMPLE 4 using the species-specific primers. Other fungal DNA species and isolates were tested for the species-specific primers ability to cross-react with them.

The S. tritici-specific primer JB446 (SEQ ID NO:12) and ITS1 (SEQ ID NO:38)(JB410) amplified a 345 bp fragment from all of the S. tritici isolates listed in EXAMPLE 1. There was no cross-reactivity with purified genomic DNA of S. nodorum, S. glycines or S. passerini. None of these other fungal species produced an amplification product with the S. tritici-specific primers.

A 448 bp fragment was amplified from all of the S. nodorum isolates listed in EXAMPLE 1 using the S. nodorum-specific primers JB433 (SEQ ID NO:7) and JB434 (SEQ ID NO:8). Similarly the S. nodorum-specific primers JB527 (SEQ ID NO:10) and JB525 (SEQ ID NO:9) amplified a 458 bp fragment from all the S. nosorum isolates listed in EXAMPLE 1. S. tritici, S. glycines and S. passerini did not produce any amplification products when assayed with the either of the S. nodorum-specific primer sets JB433 (SEQ ID NO:7) and JB434 (SEQ ID NO:8) or JB527 (SEQ ID NO:10) and JB525 (SEQ ID NO:9).

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PCRs were run using the conditions described in EXAMPLE 9, the *P. herpotrichoides*-specific primer combinations listed in EXAMPLE 8 against the other fungal DNA species and isolates listed in EXAMPLE 1.

The primer combination JB537 (SEQ ID NO:15) and JB541 (SEQ ID NO:19) produced a 413 bp fragment from the W-type P. herpotrichoides isolates only when tested against the P. herpotrichoides isolates and the following cereal pathogens: P. aestiva, C. cereale, P. sorokiniana, S. tritici and S. nodorum. The primer combiantion JB539 (SEQ ID NO:17) and

JB544 (SEQ ID NO:22) amplified a 487 bp fragment from the R-type *P. herpotrichoides* isolate only when tested against the same DNAs. The primer combination JB540 (SEQ ID NO:18) and JB542 (SEQ ID NO:20) produced a 413 bp fragment from the R-type *P. herpotrichoides* isolate only when tested against the same DNAs.

# Example 11: Sources of Pseudocercosporella herpotrichoides-infected wheat

Eyespot-infected wheat stems were received from the stage 1c fungicide screening program of Ciba Basle. Eight day old wheat plants were infected with *P. herpotrichoides* by spraying a conidial suspension (5x10<sup>5</sup> conidia/ml) in 0.2% Tween 20 on the base of the wheat stems. After inoculation, the plants were covered with plastic and incubated for one day at 20°C and 95-100% relative humidity. The plants were transferred to a growth chamber where they were incubated for four weeks at 12°C and 60% relative humidity. After this incubation, the plants were moved to a greenhouse and incubated at 18°C and 60% relative humidity. Wheat plants infected with W-type *P. herpotrichoides* strain 311 were sampled at 8-9 weeks post-infection, while those infected with the R-type strain 308 pathogen were harvested at 9-10 weeks post-infection

# Example 12: DNA extraction from wheat stems for *P. herpotrichoides* assay

DNA was extracted from wheat stems using the protocol described by Klimyuk *et al.* (The Plant Journal 3(3):493-494) with some modifications. A 2 cm wheat stem cut 0.5 cm above the basal culm was placed in 160 μl of 0.25 M NaOH and ground with a Kontes pestle until completely macerated. The sample was boiled for 30 s. 160 μl of 0.25 M HCl and 80 μl of 0.5 M Tris-Cl,pH8.0/ 0.25% v/v Nonidet P-40 were added to the sample. The sample was boiled for an additional 2 mins., then placed in an ice water bath. 1 μl of extract was used in the PCR assay.

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# Example 13: Incorporation of diagnostic assays into a quantitative colormetric assay format

The colormetric assay was performed according to Nikiforov et al. (PCR Methods and Applications 3:285-291) with the following changes:

1) 30  $\mu$ l of the R-type PCR product and 3 M NaCl/20 mM EDTA mixture were added to the capture primer well. 50  $\mu$ l of the W-type PCR product and 3 M NaCl/20 mM EDTA mixture were used in the hybridization reaction.

2) The exonuclease treatment and hybridization reaction were incubated at 37°C.

3) A 1:1000 dilution of anti-biotin horseradish peroxidase (HRP) monoclonal antibody was used.

4) After a 2 min. incubation with the O-phenylenediamine dihydrochloride (OPD) substrate, 50 µl of 3 N HCl were added to each assay well. 96-well plates were read at 492 nm and referenced at 570 nm using a conventional ELISA plate reader.

The primers listed in Table 5 were synthesized as described in EXAMPLE 5 for testing as capture primers for the colormetric assay.

Table 5: Capture Primer Design for Colormetric Assay

25 **Primer Primer Template** Primer Sequence <u>Name</u> 5'GCTGCGTTCTTCATCGATGC3' (SEQ ID NO:39) ITS2 5.8S rDNA 5'CCACTGATTTTAGAGGCCGCGAG3'(SEQ ID NO:19) JB541 W-type P. herp. 30 5'CCACTGATTTTAGAGGCCGCGAA3'(SEQ ID NO:20) JB542 R-type P. herp. 5'TGACGACTCTAAACCCTACCA3' (SEQ ID NO:66) JB538' W-type P. herp. 5'CGACGACTCTAAACCTTACCG3' (SEQ ID NO:67) JB539' R-type P. herp. 5'ATTCAAGGGTGGAGGTCTGA3' (SEQ ID NO:68) W130 W-type P. herp. 5'ATTCAAGGGTGGAGGTCTGG3' (SEQ ID NO:69) R130 R-type P. herp. 5'CTCTAAACCCTACCA3' (SEQ ID NO:70) JB538'15 W-type P. herp. 5'CTCTAAACCTTACCG3' (SEQ ID NO:71) JB539'15 R-type P. herp. 5'GTGGTCCTCTGGCAG3' (SEQ ID NO:72) JB553 R & W types

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JB554 R & W types JB555 W-type P. herp. JB556 R-type P. herp. JB561 R-type P. herp. JB562 W-type P. herp. JB559 W-type P. herp. JB550 R-type P. herp. JB557 W-type P. herp. JB558 R-type P. herp.	5'CTCAACAGCCGAAGC3' (SEQ ID NO:73) 5'GGGTGGAGGTCTGA3' (SEQ ID NO:74) 5'GGTGGAGGTCTGG3' (SEQ ID NO:75) 5'TGGAGGTCTGGACCA3' (SEQ ID NO:76) 5'TGGAGGTCTGAACCA3' (SEQ ID NO:77) 5'AGGGTGGAGGTCTGA3' (SEQ ID NO:78) 5'AGGGTGGAGGTCTGG3' (SEQ ID NO:79) 5'TTCTCCGAGAGGCCT3' (SEQ ID NO:80) 5'TTCTCCGAGAGGCCC3' (SEQ ID NO:81)
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The S. nodorum diagnostic primers JB527 (SEQ ID NO:10) and JB525 (SEQ ID NO:9) were integrated into the quantitative colormetric assay format. The primer JB527 (SEQ ID NO:10) was synthesized by Midland Certified Reagent Complany (Midland, Texas) to contain a biotin label and the 5' end to contain four internucleotidic phosphorothioate bonds. PCR amplification as described in EXAMPLE 4 using the modified JB527 (SEQ ID NO:10) and JB525 (SEQ ID NO:9) primers from healthy, low, medium, and highly S. nodorum-infected wheat produced no, low, medium and high A492 values, respectively, when assayed colormetrically using the ITS2 (SEQ ID NO:39) primer as the PCR product capture primer.

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The *P. herpotrichoides* R-type specific 5' primers, JB539 (SEQ ID NO:17) and JB540 (SEQ ID NO:18), and the *P. herpotrichoides* W-type specific 5' primer, JB537 (SEQ ID NO:15), were also modified to contain a biotin label and four internucleotidic phosphorothioate bonds. A colormetric version of the *P. herpotrichoides* R-type PCR assay was developed using the modified JB540 (SEQ ID NO:18) primer, JB542 (SEQ ID NO:20) primer and the capture primer JB539'15. The products produced from amplification from R-type infected wheat and from R-type genomic DNA using the modified JB540 (SEQ ID NO:18) primer and JB542 (SEQ ID NO:20) primer produced positive colormetric values when assayed colormetrically. Positive colormetric values were also obtained by colormetric analysis of the PCR products from amplification using the modified JB537 (SEQ ID NO:15) primer and W-type specific primer JB541 (SEQ ID NO:19)with W-type infected wheat and W-type genomic DNA when JB538'15 was used as the capture primer. Furthermore, the intensity of the colormetric signal corresponded to the fragment intensity of the PCR product as visualized on an agarose gel.

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Previously, the different Septoria species were identifiable by examination under the microscope, and the identification of the different Pseudocercosporella strains has been possible only by pathological tests. Similarly, the unambiguous identification of Mycosphaerella musicola and Mycosphaerella fijiensis has been difficult, and even the isolation of mature perithecia does not always allow accurate identification (Pons, 1990; In: Sigatoka Leaf Spot Diseases of Banana, Eds. RA Fullerton and RH Stover, International Network for the Improvement of Banana and Plantain, France). Currently immunodiagnostic kits utilizing ELISA technology are routinely used to identify Septoria tritici, Septoria nodorum, Pseudocercosporella herpotrichoides and other pathogen, but this technology lacks the accuracy, detection limit and ability to distinguish different isolates of the instant invention. In consequence, the development of a DNA test for the rapid identification of different strains of these fungi offers real advantages not only to fungal taxonomists, but also for disease management and selective fungicide use in the field.

While the present invention has been described with reference to specific embodiments thereof, it will be appreciated that numerous variations, modifications, and further embodiments are possible, and accordingly, all such variations, modifications and embodiments are to be regarded as being within the scope of the present invention.

#### **Deposits**

The following deposits were made on March 28, 1994, at Agricultural Research Service, Patent Culture Collection (NRRL), Northern Regional Research Center, 1815 North University Street, Peoria, Illinois 61604, U.S.A.:

1.	HB101 DH5d (pCRW2-1; SEQ ID NO: 3)	Accession No. NRRL B-21251
2.	HB101 DH5d (pCRW5-1; SEQ ID. NO: 47)	Accession No. NRRL B-21232
3.	E. coli DH5d (pCRSTRIT1; SEQ ID NO: 1)	Accession No. NRRL B-21233
4.	E. coli DH5d (pCRR1-21; SEQ ID NO: 4)	Accession No. NRRL B-21234
5.	E. coli DH5d (pCRSNOD31; SEQ ID NO: 2)	Accession No. NRRL B-21235

Associon No. NIDDI R-21221

#### SEQUENCE LISTING

- (1) GENERAL INFORMATION:
  - (i) APPLICANT: Ligon, James M Beck, James J
  - (ii) TITLE OF INVENTION: Detection of Fungal Pathogens Using the Polymerase Chain Reaction
  - (iii) NUMBER OF SEQUENCES: 86
  - (iv) CORRESPONDENCE ADDRESS:
    - (A) ADDRESSEE: Ciba-Geigy Corporation(B) STREET: 7 Skyline Drive

    - (C) CITY: Hawthorne
    - (D) STATE: NY
    - (E) COUNTRY: USA
    - (F) ZIP: 10532
  - (v) COMPUTER READABLE FORM:
    - (A) MEDIUM TYPE: Floppy disk
    - (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS
    - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
  - (vi) CURRENT APPLICATION DATA:
    - (A) APPLICATION NUMBER: US TBA (B) FILING DATE:

    - (C) CLASSIFICATION:
  - (vii) PRIOR APPLICATION DATA:
    - (A) APPLICATION NUMBER: US 08/233,608
    - (B) FILING DATE: 04-APR-1994
  - (viii) ATTORNEY/AGENT INFORMATION:

    - (A) NAME: Walsh, Andrea C.
      (B) REGISTRATION NUMBER: 34,988
    - (C) REFERENCE/DOCKET NUMBER: CGC 1739
    - (ix) TELECOMMUNICATION INFORMATION:
      - (A) TELEPHONE: 919-541-8666 (B) TELEFAX: 919-541-8689
- (2) INFORMATION FOR SEQ ID NO:1:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 548 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (ix) FEATURE:
    - (A) NAME/KEY: misc\_feature
    - (B) LOCATION: 1..548
    - (D) OTHER INFORMATION: /note= \*DNA sequence for the Internal Transcribed Spacer of Septoria tritici\*

(xi) SE	EQUENCE DESC	RIPTION: SE	Q ID NO:1:			
TCCGTAGGTG	AACCTGCGGA	GGGATCATTA	CCGAGCGAGG	GCCTCCGGGT	CCGACCTCCA	60
ACCCTTTGTG	AACACATCCC	GTTGCTTCGG	GGGCGACCCT	GCCGGGCGCC	CCCGGAGGAC	120
CACCAAAAAA	CACTGCATCT	CTGCGTCGGA	GTTTACGAGT	AAATCGAAAC	AAAACTTTCA	180
ACAACGGATC	TCTTGGTTCT	GGCATCGATG	AAGAACGCAG	CGAAATGCGA	TAAGTAATGT	240
GAATTGCAGA	ATTCAGTGAA	TCATCGAATC	TTTGAACGCA	CATTGCGCCC	CCTGGTATTC	300
CGGGGGCAT	GCCCGTTCGA	GCGTCATTAC	ACCACTCCAG	CCTCGCTGGG	TATTGGGCGT	360
CTTTTCGCGG	GGGATCACTC	CCCCGCGCGC	CTCAAAGTCT	CCGGCTGAGC	GGTCTCGTCT	420
					CCGTTAAATC	480
					ATATCAATAA	540
GCGGAGGA						548
GCGGAGGA						

#### (2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 583 base pairs

  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Septoria nodorum
- (ix) FEATURE:
  - (A) NAME/KEY: misc\_feature

  - (B) LOCATION: 1..583
    (D) OTHER INFORMATION: /note= DNA sequence for the Internal Transcribed Spacer of Septoria nodorum\*
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

TCCGTAGGTG	AACCTGCGGA	AGGATCATTA	CACTCAGTAG	TTTACTACTG	TAAAAGGGGC	60
TGTTAGTCTG	TATAGCGCAA	GCTGATGAGC	AGCTGGCCTC	TTTTATCCAC	CCTTGTCTTT	120
TGCGTACCCA	CGTTTCCTCG	GCAGGCTTGC	CTGCCGGTTG	GACAAATTTA	TAACCTTTTT	180
AATTTTCAAT	CAGCGTCTGA	AAAACTTAAT	AATTACAACT	TTCAACAACG	GATCTCTTGG	240
TTCTGGCATC	GATGAAGAAC	GCAGCGAAAT	GCGATAAGTA	GTGTGAATTG	CAGAATTCAG	300
TGAATCATCG	AATCTTTGAA	CGCACATTGC	GCCCCTTGGT	ATTCCATGGG	GCATGCCTGT	360
TCGAGCGTCA	TTTGTACCCT	CAAGCTCTGC	TTGGTGTTGG	GTGTTTGTCC	TCTCCCTAGT	420
GTTTGGACTC	GCCTTAAAAT	AATTGGCAGC	CAGTGTTTTG	GTATTGAAGC	GCAGCACAAG	480
TCGCGATTCG	TAACAAACAC	TTGCGTCCAC	AAGCCTTTTT	AACTTTTGAC	CTCGGATCAG	540
GTAGGGATAC	CCGCTGAACT	TAAGCATATC	AATAAGCGGA	GGA		583

121	INFORMATION	PAD.	CPA	TD	NO. 2
(2)	INFORMATION	FUR	SEO	ıυ	NO:3:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 626 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Pseudocercosporella herpotrichoides
  - (B) STRAIN: Strain R
  - (C) INDIVIDUAL ISOLATE: Variant W2-1
- (ix) FEATURE:

\*1

- (A) NAME/KEY: misc\_feature (B) LOCATION: 1..626
- (D) OTHER INFORMATION: /note= "DNA sequence for the Internal Transcribed Spacer of Pseudocercosporella herpotrichoides strain W (variant W2-1)\*
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

TCCGTAGGTG AACCTGCGGA	AGGATCATTA	ATAGAGCAAT	GAACAGACAG	CGCCCCGGGA	60
GAAATCCTGG GGGCTACCCT	ACTTGGTAGG	GTTTAGAGTC	GTCAGGCCGC	TCGGAGAAGC	120
CTGGTTCAGA CCTCCACCCT	TGAATAAATT	ACCTTTGTTG	CTTTGGCAGG	GCGCCTCGCG	- 180
CCAGCGGCTT CGGCTGTTGA	GTACCTGCCA	GAGGACCACA	ACTCTTGTTT	TTAGTGATGT	240
CTGAGTACTA TATAATAGTT	AAAACTTTCA	ACAACGGATC	TCTTGGTTCT	GGCATCGATG	300
AAGAACGCAG CGAAATGCGA	TAAGTAATGT	GAATTGCAGA	ATTCAGTGAA	TCATCGAATC	360
TTTGAACGCA CATTGCGCCC	TCTGGTATTC	CGGGGGGCAT	GCCTGTTCGA	GCGTCATTAT	420
AACCACTCAA GCTCTCGCTT	GGTATTGGGG	TTCGCGTCCT	CCCGCCTCT	AAAATCAGTG	480
GCGGTGCCTG TCGGCTCTAC	GCGTAGTAAT	ACTCCTCGCG	ATTGAGTCCG	GTAGGTTTAC	540
TTGCCAGTAA CCCCCAATTT	TTTACAGGTT	GACCTCGGAT	CAGGTAGGGA	TACCCGCTGA	600
ACTTAAGCAT ATCAATAAGC	GGAGGA				626

- (2) INFORMATION FOR SEQ ID NO:4:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 627 base pairs

    - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Pseudocercosporella herpotrichoides
    - (B) STRAIN: Strain R

		FEATURE	
ų	ix)	LEWIOKE	÷

(A) NAME/KEY: misc\_feature (B) LOCATION: 1..627

(D) OTHER INFORMATION: /note= "DNA sequence for the Internal Transcribed Spacer of Pseudocercosporella herpotrichoides Strain R\*

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

			1010100110	CCAMACACAC	CCCCCCCCC	60
TCCGTAGGTG	AACCTGCGGA	AGGATCATTA	ATAGAGCAAT	GGATAGACAG	CGCCCCGGGA	•
GAAATCCTGG	GGGCCACCCT	ACTTCGGTAA	GGTTTAGAGT	CGTCGGGCCT	CTCGGAGAAG	120
CCTGGTCCAG	ACCTCCACCC	TTGAATAAAT	TACCTTTGTT	GCTTTGGCAG	GGCGCCTCGC	180
GCCAGCGGCT	TCGGCTGTTG	AGTACCTGCC	AGAGGACCAC	AACTCTTGTT	TTTAGTGATG	240
TCTGAGTACT	ATATAATAGT	TAAAACTTTC	AACAACGGAT	CTCTTGGTTC	TGGCATCGAT	300
GAAGAACGCA	GCGAAATGCG	ATAAGTAATG	TGAATTGCAG	AATTCAGTGA	ATCATCGAAT	360
CTTTGAACGC	ACATTGCGCC	CTCTGGTATT	CCGGGGGCA	TGCCTGTTCG	AGCGTCATTA	420
TAACCACTCA	AGCTCTCGCT	TGGTATTGGG	GTTCGCGTCT	TCGCGGCCTC	TAAAATCAGT	480
GGCGGTGCCT	GTCGGCTCTA	CGCGTAGTAA	TACTCCTCGC	GATTGAGTCC	GGTAGGTTTA	540
CTTGCCAGCA	ACCCCCAATT	TTTTACAGGT	TGACCTCGGA	TCAGGTAGGG	ATACCCGCTG	600
AACTTAAGCA	TATCAATAAG	CGGAGGA				627

#### (2) INFORMATION FOR SEQ ID NO:5:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 534 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Mycosphaerella fijiensis
- (ix) FEATURE:
  - (A) NAME/KEY: misc\_feature
    (B) LOCATION: 1..534

  - (D) OTHER INFORMATION: /note= "DNA sequence for the Internal Transcribed Spacer of Mycosphaerella fijiensis"

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

TCCGTAGGTG AACCTGCGGA GGG	ATCATTA CCGAGTGAGG	GCTCACGCCC	GACCTCCAAC	60
CCTTTGTGAA CCACAACTTG TTG	CTTCGGG GGCGACCTGC	CCTCGGCGGG	CGCCCCGGA	120
GGCCGTCTAA ACACTGCATC TTT	GCGTCGG AGTTTAAAAC	AAATCGAACA	AAACTTTCAA	180
CAACGGATCT CTTGGTTCTG GCA	ATCGATGA AGAACGCAGC	GAAATGCGAT	AAGTAATGTG	240
AATTGCAGAA TTCAGTGAAT CAT	CGAATCT TTGAACGCAC	ATTGCGCCCT	TTGGTATTCC	300

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GAAGGGCATG	CCTGTTCGAG	CGTCATTTCA	CCACTCAAGC	CTGGCTTGGT	ATTGGGCGTC	360
GCGGTTCTTC	GCGCGCCTTA	AAGTCTCCGG	CTGAGCTGTC	CGTCTCTAAG	CGTTGTGGAT	420
CTTTCAATTC	GCTTCGGAGT	GCGGGTGGCC	GCGGCCGTTA	AATCTTTATT	CAAAGGTTGA	480
CCTCGGATCA	GGTAGGGATA	CCCGCTGAAC	TTAAGCATAT	CAATAAGCGG	AGGA	534

(2) INFORMATION FOR SEQ ID NO:6:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 540 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Mycosphaerella musicola
- (ix) FEATURE:
  - (A) NAME/KEY: misc\_feature
  - (B) LOCATION: 1..540
  - (D) OTHER INFORMATION: /note= "DNA sequence for the Internal Transcribed Spacer of Mycosphaerella musicola"
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

60	GACCTCCAAC	GCTCACCCCC	CCGAGTGAGG	GGGATCATTA	AACCTGCGGG	TCCGTAGGTG
120	GTCGCCGGGC	CGGCGAACTT	GCGACCCTGC	TGCTTCGGGG	CCACACCTGT	CCTTTGTGAA
180	AATCGGACAA	GTTCCAAACA	CTGCGTCGGA	CACTGCATCT	GTCTCCTTAA	GCCCCCGGAG
240	AAATGCGATA	GAACGCAGCG	CATCGATGAA	TTGGTTCTGG	AACGGATCTC	AACTTTCAAC
300	TTGCCTCCTT	TCAACGCACA	ATCGAATCTT	TCAGTGAATC	ATTGCAGAAT	AGTAATGTGA
360	TAGCTTGGTA	CACTCAAGCC	GTCATTTCAC	CTGTTCGAGC	AAGGGCATGC	TGGCATTCCG
420	CGTCTCTAAG	CTAAGCCGTC	AGTCTCCCGG	CGCGCCCCAA	CGGTGCTCCG	TTGGGCGCCG
480	AATCTTCAAA	GCGGCCGTTA	GCGGGTGGCC	GCTCCGGAGC	TTTTCAGTTC	CGTTGTGGAT
540	AAGCGGAGGA	GCATATCAAT	CTGAACTTAA	GGGATACCCG	GGATCAGGTA	GGTTGACCTC

- (2) INFORMATION FOR SEQ ID NO:7:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: Other nucleic acid
  - (A) DESCRIPTION: Oligonucleotide primer JB433
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
ACACTCAGTA GTTTACTACT	20
(2) INFORMATION FOR SEQ ID NO:8:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 17 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB434	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:	
TGTGCTGCGC TTCAATA	17
(2) INFORMATION FOR SEQ ID NO:9:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 23 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB525	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:	22
GCGACTTGTG CTGCGCTTCA ATA	23
(2) INFORMATION FOR SEQ ID NO:10:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 24 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB527	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:	

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CATTACACTC AGTAGTTTAC TACT	24
(2) INFORMATION FOR SEQ ID NO:11:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 17 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB445	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:	
CTGCGTCGGA GTTTACG	. 17
(2) INFORMATION FOR SEQ ID NO:12:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 17 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB446	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:	
CGAGGCTGGA GTGGTGT	17
(2) INFORMATION FOR SEQ ID NO:13:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 22 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	

(ii) MOLECULE TYPE: Other nucleic acid
(A) DESCRIPTION: Oligonucleotide primer JB526

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

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CCCAGCGAGG CTGGAGTGGT GT

(2) INFORMATION FOR SEQ ID NO:14:

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(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 23 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

<ul><li>(ii) MOLECULE TYPE: Other nucleic acid</li><li>(A) DESCRIPTION: Oligonucleotide primer JB536</li></ul>	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	
CTGGGGGCTA CCCTACTTGG TAG	23
(2) INFORMATION FOR SEQ ID NO:15:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB537	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:	
GGGGGCTACC CTACTTGGTA G	21
(2) INFORMATION FOR SEQ ID NO:16:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 24 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB538	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:	
ACTTGGTAGG GTTTAGAGTC GTCA	24
(2) INFORMATION FOR SEQ ID NO:17:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 24 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	

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<ul><li>(ii) MOLECULE TYPE: Other nucleic acid</li><li>(A) DESCRIPTION: Oligonucleotide primer JB539</li></ul>	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:	24
CTTCGGTAAG GTTTAGAGTC GTCG	24
(2) INFORMATION FOR SEQ ID NO:18:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 22 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB540	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:	
GGGGCCACC CTACTTCGGT AA	22
(2) INFORMATION FOR SEQ ID NO:19:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 23 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB541	•
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:	
CCACTGATTT TAGAGGCCGC GAG	23
(2) INFORMATION FOR SEQ ID NO:20:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 23 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB542	
(iii) HYPOTHETICAL: NO	

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(iv) ANTI-SENSE: NO	
·	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:	
CCACTGATTT TAGAGGCCGC GAA	23
(2) INFORMATION FOR SEQ ID NO:21:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
<ul><li>(ii) MOLECULE TYPE: Other nucleic acid</li><li>(A) DESCRIPTION: Oligonucleotide primer JB543</li></ul>	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:	
CCTGTAAAAA ATTGGGGGTT A	21
(2) INFORMATION FOR SEQ ID NO:22:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 21 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB544	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:	
CCTGTAAAAA ATTGGGGGTT G	21
(2) INFORMATION FOR SEQ ID NO:23:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 22 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB547	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:

(iii) HYPOTHETICAL: NO
 (iv) ANTI-SENSE: NO

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ATTACCGAGT GAGGGCTCAC GC

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- (2) INFORMATION FOR SEQ ID NO:24:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid
  - (A) DESCRIPTION: Oligonucleotide primer JB548
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:

GTTGCTTCGG GGGCGACCTG

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- (2) INFORMATION FOR SEQ ID NO:25:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB442
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:

TCGGGGGCGA CCTGCCG

17

- (2) INFORMATION FOR SEQ ID NO:26:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 14 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid
  - (A) DESCRIPTION: Oligonucleotide primer JB443
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:

CCGGAGGCCG TCTA

(2) INFORMATION FOR SEQ ID NO:27:

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(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 23 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB545	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:	
CCACAACGCT TAGAGACGGA CAG	23
(2) INFORMATION FOR SEQ ID NO:28:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 22 base pairs	
(B) TYPE: nucleic acid (C) STRANDEDNESS: single	
(D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB546	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:	
CACCCGCACT CCGAAGCGAA TT	22
(2) INFORMATION FOR SEQ ID NO:29:	• •
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 24 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB549	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:	
GATCCGAGGT CAACCTTTGA ATAA	24
(2) INFORMATION FOR SEQ ID NO:30:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 base pairs	

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(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB444	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:	17
GGTCAACCTT TGAATAA	•
(2) INFORMATION FOR SEQ ID NO:31:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 18 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
<ul><li>(ii) MOLECULE TYPE: Other nucleic acid</li><li>(A) DESCRIPTION: Oligonucleotide primer JB451</li></ul>	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:	
CCTTTGTGAA CCACACCT	18
(2) INFORMATION FOR SEQ ID NO:32:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 14 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB440	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:	
CTGCCGGCGA ACTT	14
(2) INFORMATION FOR SEQ ID NO:33:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 17 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	

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	(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB449	•
	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:	
-	ACCCTGCCGG CGAACTT	17
	(2) INFORMATION FOR SEQ ID NO:34:	
	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 18 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
:	(ii) MOLECULE TYPE: Other nucleic acid  (A) DESCRIPTION: Oligonucleotide primer JB448	
	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:	
	GCGACCCTGC CGGCGAAC	18
	(2) INFORMATION FOR SEQ ID NO:35:	
	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 17 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
	(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB441	
	(iii) HYPOTHETICAL: NO	
Ø.	(iv) ANTI-SENSE: NO	
Charles of the Control of the Contro	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:	
	TAGCCGGGAG ACTTTGG	17
•	(2) INFORMATION FOR SEQ ID NO:36:	
	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 16 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single	
23	(D) TOPOLOGY: linear	
	<ul><li>(ii) MOLECULE TYPE: Other nucleic acid</li><li>(A) DESCRIPTION: Oligonucleotide primer JB450</li></ul>	

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	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:	
:	TCTGCGTCGG AGTTCC	16
	(2) INFORMATION FOR SEQ ID NO:37:	
	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 18 base pairs	
:	(B) TYPE: nucleic acid (C) STRANDEDNESS: single	
10 10 10	(D) TOPOLOGY: linear	
Age.	(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer JB452	
	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
i i		
3	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:	
	CCGCGCTCCG GAGCGAAC	18
	(2) INFORMATION FOR SEQ ID NO:38:	
	(i) SEQUENCE CHARACTERISTICS:	
•	(A) LENGTH: 19 base pairs (B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
• • •	(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer ITS1	
	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
<b>Ģ</b>		
~	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:	
cg	TCCGTAGGTG AACCTGCGG	19
	(2) INFORMATION FOR SEQ ID NO:39:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 20 base pairs (B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: Other nucleic acid	
W.	(A) DESCRIPTION: Oligonucleotide primer ITS2	

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(iii) HYPOTHETICAL: NO

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:

# GCTGCGTTCT TCATCGATGC

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- (2) INFORMATION FOR SEQ ID NO:40:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer ITS3
  - (iii) HYPOTHETICAL: NO
    - (iv) ANTI-SENSE: NO
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:

### GCATCGATGA AGAACGCAGC

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- (2) INFORMATION FOR SEQ ID NO:41:
- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 20 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer ITS4
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:

## TCCTCCGCTT ATTGATATGC

- (2) INFORMATION FOR SEQ ID NO:42:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 10 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer OPB-12
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:42:	
CCTTGACGCA	10
(2) INFORMATION FOR SEQ ID NO:43:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 10 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer	OPE-6
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:43:	
AAGACCCCTC	10
(2) INFORMATION FOR SEQ ID NO:44:	
<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 10 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer	OPE-12 .
(iii) HYPOTHETICAL: NO	•
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:44:	
TTATCGCCCC	10
(2) INFORMATION FOR SEQ ID NO:45:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 10 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: Oligonucleotide primer	OPE-19
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:45:	

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- (2) INFORMATION FOR SEQ ID NO:46:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 10 base pairs
    - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid
  - (A) DESCRIPTION: Oligonucleotide primer OPE-15
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:46:

# ACGCACAACC

# (2) INFORMATION FOR SEQ ID NO:47:

- (i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 627 base pairs

  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Pseudocercosporella herpotrichoides
    - (B) STRAIN: Strain W
    - (C) INDIVIDUAL ISOLATE: Variant W5-1
  - (ix) FEATURE:
    - (A) NAME/KEY: misc\_feature
    - (B) LOCATION: 1..627
    - (D) OTHER INFORMATION: /note= "DNA sequence for the Internal Transcribed Spacer of Pseudocercosporella herpitrichoides strain W (variant W5-1) "
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:

. 60	CGCCCTGGGA	GAACAGACAG	ATAGAGCAAT	AGGATCATTA	AACCTGCGGA	TCCGTAGGTG
120	CTCGGAGAAG	CGTCAGGCCT	GGTTTAGAGT	ACTTCGGTAG	GGGCTACCCT	GAAATCCTGG
180	GGCGCCTCGC	GCTTTGGCAG	TACCTTTGTT	TTGAATAAAT	ACCTCCACCC	CCTGGTTCAG
240	TTTAGTGATG	AACTCTTGTT	AGAGGACCAC	AGTACCTGCC	TCGGCTGTTG	GCCAGCGGCT
300	TGGCATCGAT	CTCTTGGTTC	AACAACGGAT	TAAAACTTTC	ATATAATAGT	TCTGAGTACT
360	ATCATCGAAT	AATTCAGTGA	TGAATTGCAG	ATAAGTAATG	GCGAAATGCG	GAAGAACGCA
420	AGCGTCATTA	TGCCTGTTCG	CCGGGGGGCA	CTCTGGTATT	ACATTGCGCC	CTTTGAACGC
480	TAAAATCAGT	TCGCGGCCTC	GTTCGCGTCC	TGGTATTGGG	AGCTCTCGCT	TAACCACTCA

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GGCGGTGCCT CTCGGCTCTA CGCGTAGTAA TACTCCTCGC GATTGAGTCC GGTAGGTTTA	540
CTTGCCAGTA ACCCCCAATT TTTTACAGGT TGACCTCGGA TCAGGTAGGG ATACCCGCTG	600
AACTTAAGCA TATCAATAAG CGGAGGA	627
(2) INFORMATION FOR SEQ ID NO:48:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 17 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: Other nucleic acid  (A) DESCRIPTION: M13 universal -20 oligonucleotide primer	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:	
GTAAAACGAC GGCCAGT	17
(2) INFORMATION FOR SEQ ID NO:49:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 16 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
(ii) MOLECULE TYPE: Other nucleic acid (A) DESCRIPTION: M13 universal reverse oligonucleotide primer	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:49:	
AACAGCTATG ACCATG	16
(2) INFORMATION FOR SEQ ID NO:50:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 23 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid    (A) DESCRIPTION: /desc = *Oligonucleotide primer</pre>	
JB563*	

(iii) HYPOTHETICAL: NO

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(xi) SEQUENCE DESCRIPTION: SEQ 1D NO:50:	
CTTGCCTGCC GGTTGGACAA ATT	23
(2) INFORMATION FOR SEQ ID NO:51:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 25 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid     (A) DESCRIPTION: /desc = "Oligonucleotide JB564"</pre>	
(iii) HYPOTHETICAL: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:	
CTCAGTAGTT TACTACTGTA AAAGG	25
(2) INFORMATION FOR SEQ ID NO:52:	•
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 24 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid      (A) DESCRIPTION: /desc = "Oligonucleotide JB565"</pre>	
(iii) HYPOTHETICAL: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:	
CTTCTGGACG CAAGTGTTTG TTAC	24
(2) INFORMATION FOR SEQ ID NO:53:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 22 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
(iii) HYPOTHETICAL: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:	
GTTTTTAGTG GAACTTCTGA GT	22
(2) INFORMATION FOR SEQ ID NO:54:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 19 base pairs  (B) TYPE: nucleic acid  45	

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	(D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = *Oligonucleotide JB567*	
(iii)	HYPOTHETICAL: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:54:	
CGCAGGAA	CC CTAAACTCT	19
(2) INFO	RMATION FOR SEQ ID NO:55:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 12 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "Oligonucleotide primer	
JB568*	(A) DESCRIPTION: / desc - Originalization primer	
(iii)	HYPOTHETICAL: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:55:	
GCCCGCCG	CA GG	12
(2) INFO	RMATION FOR SEQ ID NO:56:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 22 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii)	MCLECULE TYPE: other nucleic acid	
JB569™	(A) DESCRIPTION: /desc = *Oligonucleotide primer	
(iii)	HYPOTHETICAL: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:56:	
RTWWTTWR	TG GAMYYTCTGA GT	22
(2) INFO	RMATION FOR SEQ ID NO:57:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 16 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "Oligonucleotide primer	

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, :	223	HYPOTHETICAL:	NO
( 1	11)	HYPOTHETICAL:	NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:57:

TATGTTGCCT CGGCGG

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- (2) INFORMATION FOR SEQ ID NO:58:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 24 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "Oligonucleotide primer JB571"
  - (iii) HYPOTHETICAL: NO
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:58:

TAACGATATG TAAATTACTA CGCT

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- (2) INFORMATION FOR SEQ ID NO:59:
- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 18 base pairs

  - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = \*Oligonucleotide primer

JB572\*

- (iii) HYPOTHETICAL: NO
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:59:

### AAGTTGGGGT TTAACGGC

- (2) INFORMATION FOR SEQ ID NO:60:
- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 14 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = \*Oligonucleotide primer JB 573\*
    - (iii) HYPOTHETICAL: NO
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:60:

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AGCGAGCCC	CG CCAC	14
(2) INFOR	RMATION FOR SEQ ID NO:61:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 22 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) JB574°	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = *Oligonucleotide primer	
(iii)	HYPOTHETICAL: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:61:	
CCATTGTG	AA CGTTACCTAT AC	22
(2) INFO	RMATION FOR SEQ ID NO:62:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 18 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: other nucleic acid	
JB575*	(A) DESCRIPTION: /desc = "Oligonucleotide primer	
(iii)	HYPOTHETICAL: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:62:	
CGACCAGA	GC GAGATGTA	18
(2) INFO	RMATION FOR SEQ ID NO:63:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: other nucleic acid	
JB576	(A) DESCRIPTION: /desc = "Oligonucleotide primer	
(iii)	HYPOTHETICAL: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:63:	
GTGAACAT	AC CTTATGTTGC C	21
(2) INFO	ORMATION FOR SEQ ID NO:64:	
(i)	SEQUENCE CHARACTERISTICS: 48	

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	<ul><li>(A) LENGTH: 16 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
(ii) JB577*	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "Oligonucleotide primer	
(iii)	HYPOTHETICAL: NO	
•		
(ix)	SEQUENCE DESCRIPTION: SEQ ID NO:64:	
GTTGCCTC	GG CGGATC	16
(2) INFO	RMATION FOR SEQ ID NO:65:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 16 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) JB578*	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "Oligonucleotide primer	
(iii)	HYPOTHETICAL: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:65:	
CCGCGACG	AT TACCAG	16
(2) INFO	RMATION FOR SEQ ID NO:66:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) JB538'	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "Oligonucleotide primer"	
(iii)	HYPOTHETICAL: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:66:	
TGACGAC	TCT AAACCCTACC A	21
(2) INF	DRMATION FOR SEQ ID NO:67:	
· (i	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii	) MOLECULE TYPE: other nucleic acid 49	

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(A) DESCRIPTION: /desc = "Ol JE539'"	igonucleotide primer
(iii) HYPOTHETICAL: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID	NO:67:
CGACGACTCT AAACCTTACC G	21
(2) INFORMATION FOR SEQ ID NO:68:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 20 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic</pre>	
(iii) HYPOTHETICAL: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID	NO:68:
ATTCAAGGGT GGAGGTCTGA	20
(2) INFORMATION FOR SEQ ID NO:69:	
<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 20 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
<pre>(ii) MOLECULE TYPE: other nucleic</pre>	acid igonucleotide primer
(iii) HYPOTHETICAL: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID	NO: 69:
ATTCAAGGGT GGAGGTCTGG	20
(2) INFORMATION FOR SEQ ID NO:70:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 15 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic      (A) DESCRIPTION: /desc = "OTITION: /desc = "OTITION"</pre>	acid ligonucleotide primer
(iii) HYPOTHETICAL: NO	

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:70:	
CTCTAAACCC TACCA	15
(2) INFORMATION FOR SEQ ID NO:71:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 15 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
(iii) HYPOTHETICAL: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:71:	
CTCTAAACCT TACCG	15
(2) INFORMATION FOR SEQ ID NO:72:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 15 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid           (A) DESCRIPTION: /desc = "Oligonucleotide primer JB553"</pre>	
(iii) HYPOTHETICAL: NO	
(xi) FEQUENCE DESCRIPTION: SEQ ID NO:72:	15
GTGGTCCTCT GGCAG	15
(2) INFORMATION FOR SEQ ID NO:73:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 15 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	·
<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
(iii) HYPOTHETICAL: NO	
	•
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:73:	
CTCAACAGCC GAAGC	15

CTCAACAGCC GAAGC

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(2) INFO	RMATION FOR SEQ ID NO:74:	
	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 14 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) JB555*	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = *Oligonucleotide primer	
(iii)	HYPOTHETICAL: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:74:	
GGGTGGAG	GT CTGA	14
	RMATION FOR SEQ ID NO:75:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 13 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: other nucleic acid	
JB556*	(A) DESCRIPTION: /desc = *Oligonucleotide primer	
(iii)	HYPOTHETICAL: NO	
GGTGGAGG' (2) INFOI (i)	SEQUENCE DESCRIPTION: SEQ ID NO:75:  TO TGG  RMATION FOR SEQ ID NO:76:  SEQUENCE CHARACTERISTICS:  (A) LENGTH: 15 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear  MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "Oligonucleotide primer"	13
	VIII DOMINIMA DA LA	
	HYPOTHETICAL: NO  SEQUENCE DESCRIPTION: SEQ ID NO:76:	
TGGAGGTC'	TG GACCA	15
	RMATION FOR SEQ ID NO:77:	
	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 15 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	

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	(D) TOPOLOGY: linear	
(ii) JB562*	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "Oligonucleotide primer	
	HYPOTHETICAL: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:77:	
TGGAGGTC'	IG AACCA	15
(2) INFO	RMATION FOR SEQ ID NO:78:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 15 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = *Oligonucleotide primer	
JB559*	(A) DESCRIPTION: /desc = *Oligonacteotide primer	
(iii)	HYPOTHETICAL: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:78:	
AGGGTGGA	GG TCTGA	15
(2) INFO	RMATION FOR SEQ ID NO:79:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 15 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii) JB560*	MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "Oligonucleotide primer	
	HYPOTHETICAL: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:79:	
AGGGTGGA	GG TCTGG	15
(2) INFO	RMATION FOR SEQ ID NO:80:	
(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 15 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: other nucleic acid	
.TD557*	(A) DESCRIPTION: /desc = "Oligonucleotide primer	

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(iii) HYPOTHETICAL: N	NU
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(xi) SEQ	QUENCE DESCRIPTION: SEQ ID NO:80:	
TTCTCCGAGA G	GCCT	15
(2) INFORMAT	rion for seq id no:81:	
( <i>1</i> (1	QUENCE CHARACTERISTICS: A) LENGTH: 15 base pairs B) TYPE: nucleic acid C) STRANDEDNESS: single D) TOPOLOGY: linear	
	LECULE TYPE: other nucleic acid	
JB558*	A) DESCRIPTION: /desc = "Oligonucleotide primer	
(iii) HYI	POTHETICAL: NO	
(xi) SE(	QUENCE DESCRIPTION: SEQ ID NO:81:	
TTCTCCGAGA (	GGCCC	15
(2) INFORMAT	TION FOR SEQ ID NO:82:	
(1 (1 (0	QUENCE CHARACTERISTICS: A) LENGTH: 504 base pairs B) TYPE: nucleic acid C) STRANDEDNESS: single D) TOPOLOGY: linear	
(ii) MOI	LECULE TYPE: DNA (genomic)	
(iii) HY	POTHETICAL: NO	
(1	A) NAME/KEY: misc_feature B) LOCATION: 1504 D) OTHER INFORMATION: /note= "DNA sequence for the transcribed spacer region of Fusarium culmorum	
(xi) SE	QUENCE DESCRIPTION: SEQ ID NO:82:	
GAGGGATCAT	TACCGAGTTT ACTRACTCCC AAACCCCTGT GAACDTACCT TATGTTGCCT	60
CGGCGGATCA	GCCCGCGCCC CGTAAAAAGG GACGGCCCGC CGCAGGAACC CTAAACTCTG	120
TTTTTAGTGG	AACTTCTGAG TATAAAAAAC AAATAAATCA AAACTTTCAA CAACGGATCT	180
CTTGGTTCTG	GCATCGATGA AGAACGCAGC AAAATGCGAT AAGTAATGTG AATTGCAGAA	240
TTCAGTGAAT	CATCGAATCT TTGAACGCAC ATTGCGCCCG CCAGTATTCT GGCGGGCATG	300
CCTGTTCGAG	CGTCATTTCA ACCCTCAAGC CCAGCTTGGT GTTGGGAGCT GCAGTCCTGC	360

420 480

TGCACTCCCC AAATACATTG GCGGTCACGT CGRAGCTTCC ATAGCGTAGT AATTTACATA

TCGTTACTGG TAATCGTCGC GGCYACGCCG TTAAACCCCA ACTTCTGAAT GTTGACCTCG

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GATCAGGTAG	GAATACCCGC	TGAA

504

(2)	INFORMATION	FOR	SEQ	ID	NO:83:
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- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 503 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO

### (ix) FEATURE:

- (A) NAME/KEY: misc\_feature
- (B) LOCATION:  $1..5\overline{03}$
- (D) OTHER INFORMATION: /note= "DNA sequence for the internal transcribed spacer region of Fusarium graminearum (fgram.con)"

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:83:

GGATCATTAC	CGAGTTTACW	SACTCCCAAA	CCCCTGTGAA	CATACCTTAT	GTTGCCTCGG	60
CGGATCAGCC	CGCGCCCCGA	AAGGGACGGC	CCGCCGCAGG	AACCCTAAAC	TCTGTTTTTA	120
GTGGAACTTC	TGAGTATAAA	AAACAAATAA	ATCAAAACTT	TCAACAACGG	ATCTCTTGGT	180
KCTGGCATCG	ATGAAGAACG	CASCRAAATG	CGATAAGTAA	TGTGWATTGC	AGAATTCAGT	240
GAATCAWCGA	ATCTTTGAAC	GCWSATTGCK	MCCRCCAGTA	TTCTGGCGGG	CATGCCTGTT	300
CGAGCGTCAT	TTCAACCCTC	AAGCCCAGVT	TGGTGTKGGG	GARYTGCAGK	CCTRYTKCAC	360
TCCCCAAATA	ARTTGGCGGT	CACGTCGAAC	TTCCATAGCG	TAGTAAGTTA	CACATCGTTA	420
CTGGTAATCG	TCGCGGCTAC	GCCGTTAAAC	CCCAACTTCT	GAATGTTGAC	CTCGGATCAG	480
GTAGGAATAC	CCGCTGAAGG	TAA				503

# (2) INFORMATION FOR SEQ ID NO:84:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 353 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO

# (ix) FEATURE:

- (A) NAME/KEY: misc\_feature
- (B) LOCATION: 1..353
- (D) OTHER INFORMATION: /note= "DNA sequence for the internal transcribed spacer region of Fusarium moniliforme (fmono.con)"

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:84:

TCCGTAGGTG AACCTGCGGA TAGGRGTCAT TASMGAGTTT ACWACTSCCA AACCCCTGTG 60

AAYATACCTT ATGTTGCSTC GGCGGATCAG CCCGCGCSCC GTARRAAGGG ACGGCCCGCC 120

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GCAGGAACCC	TAAACTCTGT	TTTTAGTGGA	ACTTCTGAGT	ATAAAAAACA	AATAAATCAA	180
AACTTTCAAC	AACGGATCTC	TTGGTTCTGG	CATCGATGAA	GAACGCAGCA	AAATGCGATA	240
AGTAATGTGA	ATTGCAGAAT	TCAGTGAATC	ATCGAATCTT	TGAACGCACA	TTGYGMCCGC	300
CAGTATTCTG	GCGGGCATGC	CTGTTCGAGC	GTCATTTCAA	CCCTCAAGCC	CAG	353

# (2) INFORMATION FOR SEQ ID NO:85:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 545 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (ix) FEATURE:
  - (A) NAME/KEY: misc\_feature
  - (B) LOCATION: 1..545
- (D) OTHER INFORMATION: /note= "DNA sequence for the internal transcribed spacer region of Microdochium nivale (mnivale.txt)\*
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:85:

GCGGATCATT .	ACAGAGTTGC	AAAACTCCCT	AAACCATTGT	GAACGTTACC	TATACCGTTG	. 60
CTTCGGCGGG	ceeccceee	GTTTACCCCC	CGGRAGYCCC	TGGKMCCCAC	CGCGGGSGCC	120
MGCCGGAGGT	CACCAAACTC	TTGATAATTT	ATGGCCTCTC	TGAGTCTTCT	GTACTGAATA	180
AGTCAAAACT	TTCAACAACG	GATCTCTTGG	TTCTGGCATC	GATGAAGAAC	GCAGCGAAAT	240
GCGATAAGTA	atgtgaattg	CAGAATTCAG	TGAATCATCG	AATCTTTGAA	CGCACATTGC	300
GCCCGCCAGC	ATTCTGGCGG	GCATGCCTGT	TCGAGCGTCA	TTTCAACCAT	CAAGCCCCCG	360
GGCTTGTGTT	GGGGACCTRC	GGCTGCCGCA	GGCCCTGAAA	AGCAGTGKCG	GGCTCGCTGT	420
CGCACCGAGM	GTAGTAGSAT	ACATCTCGCT	CTGGTCGCGC	CGCGGGTTCC	GGCCGTTAAA	480
CCACCTTTTT	AACCCAAGGT	TGACCTCGGA	TCAGGTAGGA	AGACCCGCTG	AACTTACGCA	540
TATCA						545

# (2) INFORMATION FOR SEQ ID NO:86:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 563 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (ix) FEATURE:
  - (A) NAME/KEY: misc\_feature
  - (B) LOCATION: 1..563
  - (D) OTHER INFORMATION: /note= "DNA sequence for the

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internal transcribed spacer of Septoria avenae f. sp. tricicea ATCC# 26380 (satits.con)\*

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:86:

					CT111CC1CC	60
TCCCGTAGGT	GAACCTGCGG	AAGGATCATT	ACACTCAGTA	GTTTACTACT	GTAAAGGAGG	00
CTGTTAGTCT	GTATAGCGCA	AGCTGATGAG	CAGCTAGCCT	CTTTTATCCA	CCCTTGTCTT	120
TTGCGTACCC	ACGTTTCCTC	GGCAGGCTTG	CCTGCCGATT	GGACAAACCT	ATAACCTTTT	180
TAATTTTCAA	TCAGCGTCTG	AAAAACTTAA	TAATTACAAC	TTTCAACAAC	GGATCTCTTG	240
GTTCTGGCAT	CGATGAAGAA	CGCAGCGAAA	TGCGATAAGT	AGTGTGAATT	GCAGAATTCA	300
GTGAATCATC	GAATCTTTGA	ACGCACATTG	CGCCCCTTGG	TATTCCATGG	GGCATGCCTG	360
TTCGAGCGTC	ATTTGTACCC	TCAAGCTCTG	CTTGGTGTTG	GGTGTTTGTC	CTCTCCCTAG	420
TGTTTGGACT	CGCCTTAAAA	TAATTGGCAG	CCAGTGTTTT	GGTAYTGAAG	CGCAGCACAA	480
GTCGCGATTC	TTATCAAATA	CTTGCGTCCA	CAAGCCCTTT	TTTAACTTTT	GACCTCGGAT	540
CAGGTAGGAG	ACCGCTGACT	TAA				563

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# What is claimed is:

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1. A DNA sequence encoding an Intervening Transcribed Sequence selected from the group consisting of SEQ ID NO: 1, SEQ ID NO:2, SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO: 5, SEQ ID NO: 6, SEQ ID NO: 47, SEQ ID NO: 82, SEQ ID NO: 83, and SEQ ID NO: 84, SEQ ID NO: 85, and SEQ ID NO: 86.

- 2. An oligonucleotide primer for use in amplification-based detection of a fungal Intervening Transcribed Sequence wherein said primer is derived from the DNA sequence of claim 1.
- 3. The oligonucleotide of claim 2, wherein said primer is selected from the group consisting of SEQ ID NOS: 7 to 37 and SEQ ID NOS: 50 to 65.
- 4. A pair of oligonucleotide primers for use in the amplification-based detection of a fungal Intervening Transcribed Sequence, wherein at least one primer is selected from the group consisting of SEQ ID NOS: 7 to 37 and SEQ ID NOS: 50 to 65.
- 5. The pair of oligonucleotide primers according to claim 4, wherein one primer is selected from the group consisting of SEQ ID NOS: 7 to 37 and SEQ ID NOS: 50 to 65 and the other primer is selected from the group consisting of SEQ ID NOS: 38 to 41.
  - 6. The pair of oligonucleotide primers according to claim 4, wherein said pair is selected from the group consisting of pairs of Table 4.
  - 7. The pair of oligonucleotide primers according to claim 4 wherein said pair is selected from the group consisting SEQ ID NO: 7 and SEQ ID NO: 8.
- 8. The pair of oligonucleotide primers according to claim 5, wherein said pair is selected from the group consisting of
  - (a) SEQ ID NO: 10 and SEQ ID NO: 9;
  - (b) SEQ ID NO: 12 and SEQ ID NO: 38;

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- (c) SEQ ID NO: 11 and SEQ ID NO: 41;
- (d) SEQ ID NO: 29 and SEQ ID NO: 38;
- (e) SEQ ID NO: 7 and SEQ ID NO: 41;
- (f) SEQ ID NO: 30 and SEQ ID NO: 38;
- (g) SEQ ID NO: 15 and SEQ ID NO: 19;
- (h) SEQ ID NO: 17 and SEQ ID NO: 22;
- (i) SEQ ID NO: 18 and SEQ ID NO: 20; and
- (j) SEQ ID NO: 26 and SEQ ID NO: 41.
- 9. An oligonucleotide primer for identification of a fungal pathogen, wherein said primer is selected from the group of primers consisting of SEQ ID NO: 42 to 46.
  - 10. A method for the detection of a fungal pathogen, comprising the steps of:
    - (a) isolating DNA from a plant leaf infected with a pathogen;
- 15 (b) amplifying a part of the intervening transcribed region of said pathogen using said DNA as a template in a polymerase chain reaction with a pair of primers according to claims 4 or 5; and
  - (c) visualizing said amplified part of the intervening transcribed region.
- 20 11. The method of claim 10, wherein said fungal pathogen is selected from S. nodorum, S. tritici, P. herpotrichoides, M. fijiensis, M. musicola, F. culmorum, F. graminearum, Microdochium. nivale, and F. moniliforme.
- 12. The method of claim 10, wherein said P. herpotrichoides is selected from strain W and strain R.
  - 13. A method for the detection of a fungal pathogen, comprising the steps of:
    - (a) isolating DNA from a plant leaf infected with a pathogen;
- (b) subjecting said DNA to polymerase chain reaction amplification using at least one primer according to claim 9; and
  - (c) visualizing the product or products of said polymerase chain reaction amplification.

14. A kit comprising a carrier being compartmentalized to receive in close confinement therein one or more container means, one of said container means containing the primer of claim 2.

- 5 15. A kit comprising a carrier being compartmentalized to receive in close confinement therein one or more container means, one of said container means containing the primer of claim 3.
- 16. A kit comprising a carrier being compartmentalized to receive in close confinement therein one or more container means, one of said container means containing the primers of claim 4.
  - 17. A kit comprising a carrier being compartmentalized to receive in close confinement therein one or more container means, one of said container means containing the primer of claim 5.
  - 18. A kit comprising a carrier being compartmentalized to receive in close confinement therein one or more container means, one of said container means containing the primer of claim 13.
- 19. In a quantitative colorimetric assay for the detection of a fungal pathogen comprising the steps of (a) isolating DNA from a plant leaf infected with a pathogen; (b) amplifying the DNA region of said pathogen in a polymerase chain reaction; and (c) visualizing said

amplified part of the intervening transcribed region

- wherein said improvement comprises amplifying said DNA from a part of the intervening transcribed region of said pathogen using as a template a pair of primers according to claims 4 as the diagnostic primers and visualizing said amplified part using a capture primer wherein said capture primer is selected from the group consisting of a primer of Table 5.
  - 20. The assay of claim 19 wherein said diagnostic primers are selected from SEQ ID NO.: 10 and SEQ ID NO.: 9 and the capture primer is SEQ ID NO.: 39.

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21. The assay of claim 19 wherein said diagnostic primers are selected from SEQ ID NO.: 18 and SEQ ID NO.: 20 and the capture primer is SEQ ID NO.: 71.

22. The assay of claim 19 wherein said diagnostic primers are selected from SEQ ID NO.: 15 and SEQ ID NO.: 19 and the capture primer is SEQ ID NO.: 71.

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# FIGURE

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(57) Abstract

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DNA sequences from the Internal Transcribed Spacer of the ribosomal RNA gene region are described for different spaces and strains of Septoria. Pseudocercosporella, Fusarium and Mycosphaerella. Specific primers from within these sequences are identified as being useful for the identification of the fungal isolates using PCR-based techniques.

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# Microtitration Plate Enzyme Immunoassay To Detect PCR-Amplified DNA from Candida Species ir Blood

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We developed a microtitration plate enzyme immunoassay to detect PCR-amplified DNA from Candida species. Nucleotide sequences derived from the internal transcribed spacer (ITS) region of fungal rDNA were used to develop species-specific oligonucleotide probes for Candida albicans, C. tropicalis, C. parapsilosis, and C. krusei. No cross-hybridization was detected with any other fungal, hacterial, or human DNAs tested. In contrast, a C. (Torulopsis) glabrata probe cross-reacted with Saccharomyces cerevisiae DNA but with no other DNAs tested. Genomic DNA purified from C. albicans blastoconidia suspended in blood was amplified by PCR, with fungus-specific universal primers ITS3 and ITS4. With the C. albicans-specific probe labeled with digoxigenin, a biotinylated capture probe, and streptavidin-coated microtitration plates, amplified DNA from as few as two C. albicans cells per 0.2 ml of blood could be detected by enzyme immunoassay.

Disseminated candidiasis is an important infectious complication in patients who have undergone cardiac or abdominal surgery or in patients who are severely granulocytopenic as a result of therapies for bone marrow transplantation or cancer (16, 27, 29). Antemortem diagnosis of disseminated candidiasis is difficult because the clinical presentation is usually conspecific and antibody production in immunocompromised patients can be variable, complicating the diagnosis (14). Although two or more positive blood cultures are often used to identify disseminated disease, standard blood culturing methods can require 2 to 5 days for detection and even longer for species identification (14). To shorten the time required to obtain an accurate diagnosis independent of a functioning immune system, laboratory tests have been developed to detect circulating Candida cell wall mannan (8, 21), enolase (35), or p-arabinitol (8, 14) for rapid diagnosis of disseminated candidiasis. However, the sensitivity of these tests varies among investigators and is reported to range from 22 to 100% (14).

The development of DNA-based methods for detection of Candida spp. provides an alternative and potentially more sensitive means to diagnose disseminated candidiasis. Southern blotting of nonamplified DNA targets has a detection limit of approximately 500 to 10<sup>5</sup> blastoconidia, depending on the method and probe used (6, 11). PCR technology (30) was recently adapted to amplify Candida albicans DNA, facilitating its detection (3, 7, 10, 25). However, detection of C. albicans DNA recovered from clinical specimens, even after PCR amplification, lacks sensitivity and is cumbersome for most laboratories (3, 7), particularly when DNA is recovered from blood (10, 24). Sensitivity can be improved to 10 cells per ml (15) or 3 cells per 0.1 ml (24), but this requires the use of Southern blotting coupled with radioisotopically labeled probes for detection.

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In the present study, Candida sp. 5.8S rRNA genes and the adjacent internal transcribed spacer (ITS) regions were PCR amplified by using fungus-specific universal primers ITS3 and ITS4 (36). Nonisotopic, digoxigenin-labeled oligonucleotide probes were designed on the basis of the sequence of the ITS2 region of C. albicans, C. tropicalis, C. parapsilosis, C. krusei, and C. (Torulopsis) glabrata rDNA (17). These probes were then used in a microtitration plate enzyme immunoassay (EIA) to rapidly detect and identify amplified genomic DNA from C. albicans blastoconidia introduced into blood.

# MATERIALS AND METHODS

Microorganisms and reagents. C. albicans B311 and H317 and Saccharomyces cerevisiae AB972 were obtained as previously described (19). C. tropicalis WO745, C. parapsilosis WO471, C. guilliermondii WO411, C. knusei WO701, C. (Torulopsis) glabrata WO756, and Cryptococcus neoformans var. neoformans 90-6 were obtained from lyophilized stock cultures maintained in the Mycology Culture Collection, Emerging Bacterial and Mycotic Diseases Branch, Centers for Disease Centrol and Prevention (CDC). The identity of yeast isolates was determined by carbohydrate assimilation tests performed with the AP120C kit (bioMericux Vitek, Inc., Hazelwood, Mo.), germ tube formation in serum, morphology on cornmeal agar, or for C. neoformans identification, pigment production on DL-3,4-dihydroxyphenylalanine agar. Filamentous fungal isolates Aspergillus funigatus 91-019720, A. flavus 91-019724, Histoplasma capsulatum G217B, Penicillium manneffei B-3420, and Blastomyces demaitidis 4478 were obtained from the Mycology Culture Collection, Emerging Bacterial and Mycotic Diseases Branch, CDC, Filamentous fungi were identified by colonial and nicrescopic morphology. Escherichia coli DHSa was obtained from Bethesda Research Laboratories (Gaithersburg, Md.). Staphylococcus aureus ATCC 1126 DNA was kindly provided by Tammy Bannerman (Hospital Infections Program, CDC), and Pseudomonas aeruginosa ATCC 10332 DNA was kindly provided by Arnold Steigerwalt (Division of Bacterial and Mycotic Diseases, CDC). Genomic DNA from human placenta was kindly provided by David Swan (Division of Viral and Rickettsial Diseases, CDC). All chemicals were purchased from Sigma Chemical Co., St. Louis, Mo., unless otherwise specified.

Precautions against contamination. Universal precautions suggested by Kwok and Higuchi (18) were used to eliminate possible contamination of samples. Cross-contamination by acrosols was reduced by physical separation of laboratory areas used to prepare PCRs and to analyze PCR products and by using a combination of positive-displacement pipetters and acrosol-resistant pipette tips. Other precautions included autoclaving of buffers and distilled water used for PCRs, use of fresh lots of previously aliquoted PCR reagents, and testing of appropriate negative controls, including omission of either the primer or the

DNA template during the PCR.

Purification of target DNA. Candida species and S. cerevisiae isolates were grown in 10 mi of YPG broth (1% yeast extract, 2% Bacto Peptone, 1% glucose; Difco Laboratories, Detroit, Mich.) at 37°C, and DNA was purified from lysed spheroplasts as described by Lasker et al. (20). C. neoformans genomic DNA was purified from lysed spheroplasts as described by Restrepo and Barbour (28).

Filamentous fungi were grown in 50 ml of YPG broth at 30°C for 48 h. Mycelin were harvested by filtration, washed once with sterile distilled water, and ground with a mortar and pestle in the presence of liquid nitrogen in a laminar flow biological safety cabinet. Genomic DNA was then isolated as described by Spitzer et al. (33) by repeated phenol-chloroform and chloroform extractions. Bacterial DNA was isolated by standard methods (23).

Preparation of template DNA from C. albicans blastoconidia suspended in blood. blood (10 ml) from New Zealand White rabbits (Myrtle's Rabbitry, Memphis, Tenn.) was collected from a central ear artery into lysis-centrifugation tubes (Wampole Laboratories, Cranbury, N.J.) in accordance with CDC Animal Care and Use Committee guidelines. Human blood was also tested in prelimi-

nary experiments with similar results.

To determine the sensitivity of the prototype test, blood was seeded with known numbers of Candida blastoconidia. Knowing the exact number of (viable plus nonviable) cells present was important for determining test sensitivity, since PCR methods can detect DN/s from dead, as well as viable, blastoconidia (unpublished observation). Therefore, 0, 10<sup>1</sup>, 10<sup>2</sup>, 10<sup>3</sup>, or 10<sup>4</sup> C. albicans 18311 blastoconidia which had been enumerated microscopically with a hemacytometer were seeded into 1 ml of collected blood. Erythrocytes and leukocytes were lysed by adding 0.8 ml of TE buffer (10 mM Tris, 1 mM EDTA, pH 8.0) containing 0.05% (wt/vol) proteinase K (15 U/mg) and 0.05% (vol/vol) Tween 20 detergent to 0.2 ml of seeded blood. After incubation at 55°C for 30 min, blastoconidia were centrifuged at 10,000 × g for 8 min at 20°C in a Beckman Microfuge and washed twice with TE buffer containing 0.5% Tween 20 and twice with SE solution (1 M sorbitol-0.1 M EDTA, pH 7.5).

Spheroplasts were prepared by adding 0.5 ml of SE solution containing 0.1% (wavel) Zymolyase-100T (100,000 U/g; Seikagaku Corp., Tokyo, Japan) and 1% (volvol) 2-mercaptoethanol to the cell pellets. After incubation at 37°C for 30 min and an additional 30 min of incubation at 25°C on a rocker platform (20 cycles per min), spheroplasts were centrifuged at 7,000  $\times$  g for 5 min at 20°C and washed twice with SE solution. Spheroplasts were lysed in 0.4 ml of TE buffer containing 0.05% (wt/vol) proteinase K (15 U/mg) and 0.5% (vol/vol) Tween 20, incubated at 55°C for 1 h, and then heated at 95°C for 10 min to inactivate the proteinase K. Nucleic acids were extracted by adding an equal volume of TEsaturated phenol-chloroform (1 volume of TE-saturated phenol to 1 volume of chloroform) and vortex mixing for 1 min. The emulsion was centrifuged at 10,000 × g for 5 min, and the aqueous phase was extracted by adding an equal volume of chloroform. One-tenth volume of 3.0 M sodium acetate buffer (pH 5.2) was then added to the resultant aqueous phase, and DNA was precipitated by adding 2 volumes of cold isopropanol and placing the samples at -20°C for 1 h. Precipitated nucleic acids were then collected by centrifugation at  $10,000 \times g$  for 10 min at 4°C. One milliliter of 70% ice-cold ethanol was added to wash the pellet, and samples were centrifuged for an additional 5 min at  $10,000 \times g$ . Nucleic acids were then vacuum dried and resuspended in 30 µl of distilled water or TE buffer.

Oligonucleotide synthesis of primers and probes. Synthetic oligodeoxyribonucleotides were prepared by B-cyanocthyl phosphoramidite chemistry with a 380B automated DNA synthesizer (Applied Biosystems, Foster City, Calif.). The oligonucleotide primer pair ITS3 and ITS4 was previously shown to amplify fungal 5.8S rDNA and the adjacent ITS region (36). Oligonucleotide probes specific for Candida species were prepared from sequences of the rDNA ITS2 region (17, 22, 36). Oligonucleotide probes CA, CT, CP, CK, and CG were designed to detect C. albicans, C. tropicalis, C. parapsilosis, C. krusei, and C. (Torulopsis) glabrata, respectively. For probes CA, CT, and CP, nonhomologous regions of ITS2 were derived from GenBank entries L07796, L11349, and L11352, respectively. Probes CG and CK were developed on the basis of data

previously reported (17).

Oligonucleotide probes were initially synthesized with a 5'-terminal amine group (Aminolink 2; ABI, Foster City, Calif.). Anino-linked oligonucleotides were mixed with a 10-fold molar excess of digoxigenin-3-O-methylcarbonyl-caminocaproie acid N-hydroxysuccinimide ester (Boehringer Mannheim, Indianapolis, Ind.) in 0.1 M sodium carbonate buffer, pH 9.0. After overnight incubation at ambient temperature, the digoxigenin-conjugated oligonucleotides were purified by reverse-phase high-pressure liquid chromatography (2).

The 5.8S rDNA consensus oligonucleotide probe BP was labeled with biotin by incorporating dimethoxytrityl-biotin-carbon 6-phosphoramidite (Cambridge Research Biochemicals, Inc., Wilmington, Del.) at the 5' end during standard synthesis on a 380B DNA synthesizer. The dimethoxytrityl group was retained on the biotinylated oligonucleotide to facilitate purification by reverse-phase high-pressure liquid chromatography (2). ITS4P (5'-end-protected 3' primer) was synthesized as previously described (34).

PCR conditions. A Gene Amp DNA amplification reagent kit (Perkin-Elmer Corp., Norwalk, Conn.) was used for PCR amplification of genomic DNA. The reaction mixture (50 µl) contained 50 mM KCl; 1.5 mM MgCl<sub>2</sub>; 0.001% gelatin; 10 mM Tris-HCl (pH 8.3); 50 μM each dATP, dCTP, dGTP, and dTTP; 0.1 μM each primer; 5 nl of template DNA, and 1.25 U of Taq DNA polymerase. A primer concentration of 0.1 µM was found by titration to be optimal for DNA

amplification with minimal primer dimerization (unpublished observations). Samples were overlaid with 30 µl of mineral oil prior to PCR amplification in a Perkin Eimer Cetus DMA thermal cycler. Taq polymerase was added after the thermal cycler reached 94°C and before initiation of temperature cycling. PCR amplification was determined to be optimum after 30 cycles of denaturation at 95°C for 1 min, primer annealing at 55°C for 1 min, and extension at 72°C for 1.5 min. After the last cycle, final DNA extension was performed at 72°C for 5 min.

Agarose gel electrophoresis. Electrophoresis was conducted in TBE buffer (0.089 M Tris, 0.089 M boric acid, 0.02 M EDTA, pH 8.4) at 80 V for 1 to 2 h on gels composed of 1% (wt/vol) agarose (International Technologies, Inc. New Haven, Conn.) and 1% (wt/vol) NuSieve (FMC Bioproducts, Rockland, Maine).

Microtitration plate hybridization assay. PCR-amplified DNA was hybridized to digoxigenin- and biotin-labeled oligonucleotide probes and detected in an ELA by capture with streptavidin-coated microtitration plates (Immulon 2; Dynatech Laboratories, Inc., Chantilly, Va.) (9). To facilitate hybridization, single-stranded DNA was prepared from double-stranded PCR products by either exonuclease digestion of the non-phosphothioate-protected strand (9) or heat denaturation as described below.

Ten microliters of the PCR product was supplemented with dithiothreitol to ! mM and digested with 0.4 U of T7 gene 6 exonuclease (United States Biochemical, Cleveland, Ohio) per µl for 15 min at 37°C. The digested product was then heated at 75°C for 15 min to inactivate the exonuclease. Alternatively, 10 µl of the PCR product was heated at 95°C for 5 min and then immediately cooled on ice. The single-stranded PCR product obtained by either exonuclease digestion or heating was added to 0.2 ml of hybridization solution (4× SSC (saline sodium citrate buffer; 0.6 M NaCl, 0.06 M trisodium citrate, pH 7.0] containing 20 mM N-2-hydroxyethylpiperazine-N'-2-ethanesulfonic acid [HEPES], 2 mM EDTA, and 0.15% [vol/vol] Tween 20) supplemented with 50 ng each of biotin- and

digoxigenin-labeled probes per ml.

Hybridization reactions were performed in 0.5-ml Eppendorf tubes at 37°C for 1 h. After hybridization, 100 μl of each sample was added to duplicate wells of the streptavidin-coated microtitration plate (9), and the plate was incubated at ambient temperature for 1 h with shaking (Minishaker, manufactured for Dynatech by CLTI, Middletown, N.Y.). After washing with potassium phosphate-buffered saline containing 0.05% Tween 20, 100 µl of peroxidase-conjugated. anti-digoxigenin Fab fragment (Boehringer Mannheim) diluted 1:2,000 in hybridization buffer was added per well. Plates were subsequently washed six times with potassium phosphate-buffered saline-0.05% Tween 20. Each well received 100 µl of a mixture of 1 volume of 3, 3', 5, 5'-tetramethylbenzidine peroxidase substrate (Kirkegaard & Perry Laboratories, Inc., Gaithersburg, Md.) and 1 volume of peroxidase solution (Kirkegaard & Perry). The plates were then placed at ambient temperature for 15 min, and the  $A_{650}$  of each well was determined with a microtitration plate reader (UV Max; Molecular Devices, Inc., Menlo Park, Calif.). The absorbance of a reagent blank, in which the test sample was replaced with distilled water, was subtracted from each test sample,

Statistical analyses. Student's t test was used to determine significant differences between means plus or minus the standard deviation from the mean. P

values of <0.05 were considered significant.

# RESULTS

PCR amplification of fungal rDNA with universal fungal primers ITS3 and ITS4. All of the oligodeoxyribonucleotide primers and oligonucleotide probes used in this study are described in Table 1. One nanogram of genomic DNA from each isolate was amplified by PCR with the universal fungal primer pair ITS3 and ITS4. Use of this primer pair resulted in amplification of DNAs from all of the fungi examined, including C. albicans, C. tropicalis, C. parapsilosis, C. guilliermondii, C. krusei, C. (Torulopsis) glabrata, S. cerevisiae, C. neoformans, A. fumigatus, A. flavus, P. marneffei, B. dermatitidis, and H. capsulatum (Fig. 1). No amplicon was detected with DNA isolated from S. aureus, E. coli, P. aeruginosa, or a human placental cell line. The following amplicon sizes were obtained for Candida species: 330 bp for C. albicans, 325 bp for C. tropicalis, 310 bp for C. parapsilosis, 260 bp for C. guilliermondii, 335 bp for C. krusei, and 410 bp for C. (Torulopsis) glabrata (Fig. 1). Fungi tested not belonging to a Candida species yielded amplicons with sizes ranging from 340 to 410 bp (Fig. 1).

Comparison of EIA sensitivity for detection of a PCR product by heat denaturation versus exonuclease digestion. Two methods for the production of single-stranded capture DNA to coat microtitration plates were compared. One method used heat to denature DNA into single strands, whereas the other